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- 71) Applicant: TERUMO Kabushiki Kaisha 44-1 Hatagaya 2-chome Shibuya-ku Tokyo (JP)
- (72) Inventor: Sudo, Tadashi c/o Terumo K.K., 1500 Inokuchi, Nakai-machi Ashigarakami-gun, Kanagawa-ken (JP) inventor: Harada, Kazumichi c/o Terumo K.K., 1500 Inokuchi, Nakai-machi Ashigarakami-gun, Kanagawa-ken (JP) Inventor: Hirahara, Ichiro c/o Terumo K.K., 1500 Inokuchi, Nakai-machi Ashigarakami-gun, Kanagawa-ken (JP) Inventor: Adachi, Masami c/o Terumo K.K., 1500 Inokuchi, Nakai-machi Ashigarakami-gun, Kanagawa-ken (JP)
- (74) Representative: Gillard, Marie-Louise et al Cabinet Beau de Loménie 158, rue de l'Université F-75340 Paris Cédex 07 (FR)
- (54) Vascular endothelial cells growth factor.
- A novel protein of human origin produced by a human ovarian tumor established cell line HUOCA-II or HUOCA-III, which has a molecular weight, when determined by SDS-polyacrylamide gel electrophoresis, of from 72,000 to 80,000 daltons under a non-reducing condition or from 79,000 to 85,000 daltons under a reducing condition, which contains an amino acid sequence represented by the Sequence ID No. 4 deduced from the DNA sequence represented by the Sequence ID No. 5, and which enhances growth of vascular endothelial cells but does not activate growth of smooth muscle cells, fibroblasts and hepatocytes and also does not enhance or inhibit growth of HeLa cells. This invention also provides a process for the production of the protein.

FIELD OF THE INVENTION

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This invention relates to a novel protein of human origin and its production process. Particularly, it relates to a novel proteinous angiogenic factor of human origin, which enhances the growth of vascular endothelial cells but does not activate the growth of other cells such as smooth muscle cells, fibroblasts, hepatocytes and the like, and to a process for the production thereof.

BACKGROUND OF THE INVENTION

Principal cells which constitute a blood vessel are vascular endothelial cells of tunica intima, smooth muscle cells of tunica media and fibroblasts of tunica externa. In addition, peripherally existing capillary blood vessels are composed solely of vascular endothelial cells. Though the mechanism of new formation of blood vessels, or angiogenesis, has not yet been elucidated in full details, it is considered that the angiogenesis starts firstly with dissolution of the blood vessel wall matrix and subsequent growth and migration of vascular endothelial cells.

Angiogenesis can be found during the prenatal period when new tissues and blood vessels are formed and at the time of the occurrence of physiological phenomena in the adult body such as periodical development of uterine endometrium and lutenization in ovaries, as well as under pathologic conditions such as chronic inflammation, wound healing and the like. New formation of blood vessels can also be found at the time of the growth of tumor cells. Endothelial cells which cover the inner wall of blood vessels are possessed of many physiological functions such as maintenance of anti-thrombotic activity, regulation of matter permeation, regulation of blood pressure and the like. In a patient suffering from a blood vessel-related disease such as arteriosclerosis, myocardial infarction or the like, abnormality can be found in these blood vessel-constituting cells.

A number of angiogenic factors have been found in the *in vivo* experimental systems for the formation of new blood vessels, such as an experiment in which chick chorio-allantoic membrane is used. For example, generally known proteinous angiogenic factors include basic fibroblast growth factor (bFGF), epidermal growth factor (EGF), platelet-derived growth factor (PDGF), transforming growth factor (TGF) and the like.

Though these prior art angiogenic factors having the ability to enhance formation of new blood vessels are possessed of the activity to enhance growth of vascular endothelial cells, these factors also strongly activate growth of other cells. For example, bFGF activates growth of various cells such as fibroblasts, smooth muscle cells, epidermal cells and the like. In consequence, each of these prior art angiogenic factors having a broad range of growth enhancing effects on various types of cells enhances not only the formation of new blood vessels but also the growth of other cells at the same time. In other words, these prior art factors have a problem of causing secondary reactions when used because of their inability to selectively enhance formation of new blood vessels.

Accordingly, the present invention contemplates overcoming the aforementioned problems involved in the prior art and, as the results, providing a purified angiogenic factor which enhances growth of vascular endothelial cells but does not or hardly activate growth of other cells such as smooth muscle cells, fibroblasts, hepatocytes and the like. The present invention also contemplates developing side effect-free pharmaceutical preparations and medical devices based on such a purified angiogenesis factor.

With the aim of accomplishing these objects, the inventors of the present invention have conducted intensive .studies and found that products of human ovarian tumor established cell lines, HUOCA-II and HUOCA-III, were able to enhance growth of vascular endothelial cells selectively. The results have been disclosed in Japanese Patent Application Kokai Nos. 2-261375, 2262523 and 3-84000.

Thereafter, the present inventors have carried out studies on the purification of the aforementioned products of HUOCA-II and HUOCA-III cell lines from their serum-free culture supernatants, making use of specific purification techniques, and have succeeded in obtaining a highly purified specific protein having the aforementioned desirable properties, that is, having a strong activity to enhance growth of vascular endothelial cells but with no activity to activate growth of other cells such as smooth muscle cells, fibroblasts, hepatocytes and the like.

By further continuing the studies, a total RNA was isolated from the HUOCA-II or HUOCA-III cells and its cDNA was cloned. Thereafter, the DNA sequence of the cDNA was determined and its corresponding amino acid sequence was deduced, thereby succeeding in obtaining the novel protein of the present invention.

SUMMARY OF THE INVENTION

According to a first aspect of the present invention, there is provided a single chain protein produced by

HUOCA-II or HUOCA-III, which has the following properties of:

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- (1) having a molecular weight, when determined by SDS polyacrylamide gel electrophoresis, of from 72,000 to 80,000 daltons under a non-reducing condition or from 79,000 to 85,000 daltons under a reducing condition;
- (2) containing three peptide chains, respectively represented by the Sequence ID Nos. 1, 2 and 3 as attached hereto (in the Sequence ID No. 3, "Xaa" means an unidentified amino acid residue), in one molecule;
- (3) having an activity to enhance the growth of vascular endothelial cells;
- (4) having no activity to enhance the growth of ubroblasts; vascular smooth muscle cells and hepatocytes;
- (5) having no activity to enhance or inhibit the browth of HeLa cells; and
- (6) having an activity to enhance formation of new blood vessels.

According to a second aspect of the present invention, there is provided a protein of human origin which contains an amino acid sequence or a portion of the amino acid sequence represented by the Sequence ID No. 4 attached hereto that has been identified by isolating a corresponding total RNA molecule from HUOCA-II or HUOCA-III cells, cloning a cDNA corresponding to the proteins, determining the DNA sequence of the cDNA and deducing an amino acid sequence from the DNA sequence.

According to a third aspect of the present invention, there is provided a process for the production of a protein of human origin according to the first or second aspect of the present invention, which comprises purifying a serum-free culture supernatant of a human ovarian tumor cell or established cell line thereof, especially HUOCA-II or HUOCA-III, by an optional combination of purification techniques including (a) cation exchange chromatography, (b) heparin affinity chromatography, (c) heparin affinity high performance liquid chromatography and (d) reverse phase high performance liquid chromatography, or which comprises the steps of (i) preparing a DNA fragment containing a nucleotide sequence which encodes the protein or a portion of the protein shown in the Sequence ID No. 4 attached hereto, (ii) obtaining a transformant by transforming cells of a host with the DNA fragment prepared in the above step (i) or with a vector containing the DNA fragment and (iii) culturing the transformant obtained in the above step (ii) to allow the transformant to produce the protein of the Sequence ID No. 4, or a portion of the protein, subsequently recovering the protein from resulting culture mixture.

According to a fourth aspect of the present invention, there is provided a pharmaceutical preparation which contains the protein or a portion of the protein of the first and/or second aspect of the present invention as an active ingredient

According to a fifth aspect of the present invention, there is provided a DNA fragment or cDNA-fragment which contains a nucleotide sequence or a portion of the nucleotide sequence represented by the Sequence ID No. 5 attached hereto wherein at least one base may be substituted based on the degeneracy of genetic code.

According to a sixth aspect of the present invention, there is provided an expression vector containing the DNA fragment, as well as a transformant transformed with the DNA fragment or the expression vector.

Other objects and advantages of the present invention will be made apparent as the description progresses.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is a graph showing the absorbance, measured at a wave length of 280 nm, of each eluate fraction resulting from the treatment of an HUOCA-III serum-free culture supernatant with cation exchange chromatography.

Fig. 2 is a graph showing the results of the measurement of activities in the eluate fractions obtained in Fig. 1 to enhance the growth of vascular endothelial cells.

Fig. 3 is a graph showing the absorbance, measured at a wave length of 280 nm, of each eluate fraction resulting from a heparin affinity chromatographic treatment of the active fractions of the cation exchange chromatography eluates having the vascular endothelial cell growth-enhancing activity.

Fig. 4 is a graph showing the results of the measurement of activities in the eluate fractions obtained in Fig. 3 to enhance the growth of vascular endothelial cells.

Fig. 5 is a graph showing the absorbance, measured at a wave length of 215 nm, of each eluate fraction resulting from a heparin affinity high performance liquid chromatographic treatment of the active fractions of the heparin affinity chromatography eluates having the vascular endothelial cell growth-enhancing activity.

Fig. 6 is a graph showing the results of the measurement of activities in the eluate fractions obtained in Fig. 5 to enhance growth of vascular endothelial cells.

Fig. 7 is a graph showing the absorbance, measured at a wave length of 215 nm, of each eluate fraction

resulting from a reverse phase high performance liquid chromatographic treatment of the active fractions of the heparin affinity high performance liquid chromatography eluates having the vascular endothelial cell growth-enhancing activity.

Fig. 8 is a graph showing the results of the measurement of activities in the eluate fractions obtained in Fig. 7 to enhance the growth of vascular endothelial cells.

Fig. 9 is a graph showing an SDS polyacrylamide gel electrophoresis pattern of a highly purified product (glycoprotein) obtained in Example 1 of the present invention.

Fig. 10 is a graph showing results of the measurement of the vascular endothelial cell growth-enhancing activity of the highly purified product eluted from each cut portion of the electrophoresis gel of Fig. 9.

Fig. 11 is a graph showing an SDS-polyacrylamide gel electrophoresis pattern of an N-glycanase-treated product of the highly purified product (glycoprotein) obtained in Example 1 of the present invention.

Fig. 12 represents the nucleotide sequence of the mRNA from which the cDNA obtained in Example 1 step (B) is translated and the corresponding amino acid sequence deduced from the nucleotide sequence.

DETAILED DESCRIPTION OF THE INVENTION

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Firstly, a first and a second aspects of the present invention are described in detail.

The gist of the first aspect of the present invention resides in a single chain protein produced by HUOCA-III or HUOCA-III, which has the following properties of:

- (1) having a molecular weight, when determined by SDS polyacrylamide gel electrophoresis, of from 72,000 to 80,000 daltons under a non-reducing condition or from 79,000 to 85,000 daltons under a reducing condition;
- (2) containing three peptide chains, respectively represented by the Sequence ID Nos. 1, 2 and 3 as attached hereto (in the Sequence ID No. 3, "Xaa" means an unidentified amino acid residue), in one molecule;
- (3) having an activity to enhance the growth of vascular endothelial cells;
- (4) having no activity to enhance the growth of fibroblasts, vascular smooth muscle cells and hepatocytes;
- (5) having no activity to enhance or inhibit the growth of HeLa cells; and
- (6) having an activity to enhance the formation of new blood vessels.

The gist of the second aspect of the present invention resides in a protein of human origin which contains an amino acid sequence or a portion of the sequence represented by the Sequence ID No. 4 attached hereto that has been identified by isolating a corresponding mRNA molecule from HUOCA-II or HUOCA-III cells, cloning a gene corresponding to the mRNA, determining the DNA sequence of the gene and deducing an amino acid sequence from the DNA sequence.

The human ovarian tumor established cell lines HUOCA-II and HUOCA-III have been deposited by the present inventors on March 1, 1989, in Fermentation Research Institute, Agency of Industrial Science and Technology, and have been assigned the designations as FERM BP-2310 and FERM BP-2311. Though culturing of the HUOCA-III and HUOCA-IIII and preparation of their serum-free culture supernatants may be carried out in the usual way, these techniques are disclosed in detail by the present inventors in Japanese Patent Application Kokai Nos. 2-261375, 2-262523 and 3-84000.

The protein of the present invention comprises a single chain protein molecule, and the single chain protein contains three peptide chains respectively represented by the Sequence ID Nos. 1, 2 and 3 as attached hereto.

The protein of the present invention may be prepared from a serum-free culture supernatant of the human ovarian tumor established cell line, HUOCA-II or HUOCA-III, by subjecting the supernatant to a series of purification steps including (a) cation exchange chromatography, (b) heparin affinity chromatography, (c) heparin affinity high-performance liquid chromatography and (d) reverse-phase high-performance liquid chromatography. Preferably, it may be prepared in accordance with the following illustrative steps (i) to (iv).

Preparation of protein

(i) A serum-free culture supernatant of HUOCA-II or HUOCA-III is adsorbed on to a cation exchange resin packed in a column. In this instance, the cation exchange resin may be either strongly ionic or weakly ionic, but the use of S-Sepharose® (trademark of Pharmacia) is particularly preferred. The thus adsorbed portion onto a cation exchange resin in the column is washed with an appropriate buffer solution and then subjected to a linear gradient elution using two buffer solutions respectively containing 150 mM NaCl and 2 M NaCl to collect active fractions showing the activity to enhance the growth of vascular endothelial cells [step (a)].

(ii) The active fractions obtained in the above step (i) are pooled and diluted by a factor of 2 to 3 with the

same buffer solution containing 150 mM of NaCl. The thus diluted sample is applied to a heparin-Sepharose column, washed with the same buffer solution containing 0.5 M NaCl and then subjected to a linear gradient elution using two buffer solutions respectively containing 0.5 M NaCl and 2 M NaCl to collect active fractions showing the activity to enhance the growth of vascular endothelial cells [step (b)].

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- (iii) The active fractions obtained in the above step (ii) are diluted in the same manner, applied to a heparin column for high performance liquid chromatography use and then subjected to elution in the same manner to collect active fractions showing the activity to enhance the growth of vascular endothelial cells [step (c)].
- (iv) The active fractions obtained in the above step (iii) are applied to a column for reverse-phase high-performance liquid chromatography use to obtain a purified product (protein) having the activity to enhance the growth of vascular endothelial cells [step (d)].

Any usually used buffer solution such as a phosphate buffer or the like may be used in the above glycoprotein preparation steps, and Sepharose or any other general purpose carrier may be used as a carrier of heparin.

The thus purified product has been identified as a glycoprotein, namely a sugar chain-attached protein molecule, on the basis of the facts that (1), when the purified product was allowed to react with a sugar chain-hydrolyzing enzyme N-glycanase and the resulting product was analyzed by 0.1% SDS-containing 10% polyacrylamide gel electrophoresis, the electrophoresis pattern of the thus treated product showed a decreased molecular weight level due to the digestion of sugar chains and (2) the purified product showed an affinity for concanavalin A.

In addition, the protein portion of the glycoprotein of the present invention was identified as a single chain protein molecule, because the purified product showed a single band when analyzed by 0.1% SDS-containing 10% polyacrylamide gel electrophoresis under reducing conditions.

Though the amino acid sequence of the protein portion of the thus obtained glycoprotein could be determined by any usually used means, the following illustrative steps (1) to (3) were employed herein in that order.

Determination of amino acid sequence

(1) Reductive carboxymethylation

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The sample purified and isolated in the aforementioned step (iv) by reverse-phase high-performance liquid chromatography was concentrated using a concentrator and eluted with an eluting solution consisting of 8 M urea, 0.5 M Tris-HCl pH 8.0 and 1 mM EDTA. To this was added dithiothreitol to a final concentration of 20 mM. After nitrogen gas flush, the reduction reaction was carried out in the dark for 2 hours at room temperature. Thereafter, monoiodoacetic acid was added to the resulting reaction mixture to a final concentration of 20 mM, and the alkylation reaction was carried out in the dark for 30 minutes at room temperature.

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(2) Digestion with lysyl endopeptidase

The reductive alkylation product obtained in the above step (1) was mixed with 2-mercaptoethanol, followed by the addition of 0.1 N NaOH to adjust the mixture to pH 8.5. Lysyl endopeptidase (Wako Pure Chemical Industries, Ltd.) was added in a 1:10 (w/w) ratio to the thus prepared substrate to carry out the enzymatic hydrolysis reaction at 37°C for 4 hours.

(3) Fractionation of peptide fragments and determination of the amino acid sequence

The peptide fragments mixture obtained in the above step (2) were separated by reverse-phase high-performance chromatography using an RP300 column (Applied Biosystems, Inc.). The elution was carried out by linear concentration gradient of acetonitrile from 0% to 60% in the presence of 0.1% TFA. The thus obtained peptide fragments by the elution treatment were subjected to Edman degradation using a gas phase sequencer (Model 477A; Applied Biosystems, Inc.), and the resulting PTH-amino acids were identified using a high-performance liquid chromatography for PTH-amino acid identification use (Model 120A; Applied Biosystems, Inc.). As the results, if was found that the protein portion of the glycoprotein of the present invention contained three peptide chains respectively represented by the Sequence ID Nos. 1, 2 and 3.

Determination of the complete DNA sequence by PCR

The amino acid sequence determined in the above step (3) coincided well with that of human hepatocyte

growth factor (hHGF). With regard to hHGF, its cDNA sequence has been reported by Nakamura (*Nature*, vol.342, pp.440 - 443, 1989) and Miyazawa (*Biochemical and Biophysical Research Communication*, vol.163, pp.967 - 973, 1989).

Since several cDNA nucleotide sequences have been reported on the hHGF family, primers for PCR use were prepared using a DNA synthesizer based on the common sequences in the 5' and 3' non-translation regions of these known nucleotide sequences. That is, primers were synthesized based on a region including 47 to 82 position bases (5' primer) counting in upstream direction from the 5' end of the translation region (translation initiation point) and another region including 1 to 37 position bases (3' primer) counting in downstream direction from the 3' end.

The total RNA sample was prepared from the human ovarian tumor cell line HUOCA-III by means of an SDS-phenol method. Using the thus prepared total RNA as a template, cDNA synthesis was carried out making use of M-MLV reverse transcriptase. The thus synthesized cDNA was subjected to PCR and the resulting PCR product was applied to agarose gel electrophoresis to find a DNA fragment having a size of about 2.3 kb. Since the open reading frame of the HGF family so far reported has a size of about 2.3 kb, this DNA fragment was considered to be a cDNA molecule coding for the HUOCA-III-originated novel protein of the present invention. In consequence, this DNA fragment was purified from the agarose gel, inserted into the pUC18 plasmid vector and then transformed into Escherichia coli JM109. Some of the thus obtained dones were examined making use of the dideoxy method to determine their nucleotide sequences. By correcting reading errors at the time of the PCR study, a nucleotide sequence corresponding to the novel protein of HUOCA-III origin was determined. The thus determined nucleotide sequence is shown in the Sequence ID No. 5 attached hereto, and an amino acid sequence deduced from the nucleotide sequence in the Sequence ID No. 4

Measurement of molecular weight by SDS-polyacrylamide gel electrophoresis

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Electrophoresis was carried out using a 10% polyacrylamide gel in accordance with the procedure of Lammeli et al. (Nature, vol.277, pp.680 - 685, 1970). The resulting gel was fixed by treating it with 50% ethanol and 40% acetic acid for 30 minutes, washed with 10% ethanol and 5% acetic acid and then subjected to silver staining. The protein of the present invention was stained as a single band, and its molecular weight was estimated to be about 72,000 to 80,000 daltons based on its relative mobility. In addition, another electrophoresis was carried out under a reducing condition by adding 2-mercaptoethanol to the sample to a concentration of 5% and treating the mixture at 95°C for 10 minutes, followed by the same procedure as the case of the above non-reducing condition. Under the reducing condition, the molecular weight of the protein of the present invention was estimated to be about 79,000 to 85,000 daltons.

Next, a third aspect of the present invention is described in the following.

The gist of the third aspect of the present invention resides in a process for the production of the protein of the first or second aspect of the present invention.

Firstly, a culture mixture containing the protein of the first or second aspect of the present invention is obtained.

The single chain protein of the first aspect of the present invention is obtained by recovering it from a serum-free culture supernatant of the human ovarian tumor cell line, HUOCA-II or HUOCA-III

The novel protein of the second aspect of the present invention is obtained by preparing a DNA fragment containing a nucleotide sequence which encodes the novel protein represented by the amino acid sequence or a portion of the sequence shown in the Sequence ID No. 4, preferably the DNA fragment or a portion of the DNA fragment represented by the Sequence ID No. 5, transforming appropriate host cells with the thus ligated fragment directly or indirectly using a proper expression vector, culturing the thus obtained transformant and then recovering the novel protein of the Sequence ID No. 4 from the resulting culture mixture.

The recovering step may be effected, though not particularly limited, by purifying the novel protein by means of (a) cation exchange chromatography, (b) heparin affinity chromatography, (c) heparin affinity high-performance liquid chromatography and (d) reverse-phase high-performance liquid chromatography, in any optional combination or order.

According to a fourth aspect of the present invention, there is provided a pharmaceutical preparation which contains the protein of the first and/or second aspect of the present invention as an active ingredient.

The pharmaceutical preparation may be applied to various dosage forms such as tablets, sugar coated tablets, powders, capsules, granules, suspensions, emulsions, parenteral solutions, external preparations, ointments and the like, using the preparation alone or together with other necessary ingredients in combination with appropriate carriers, fillers and the like.

The protein of the present invention is possessed of a function to enhance vascular endothelial cell growth in human and various animals, but does not enhance the growth of fibroblasts, vascular smooth muscle cells

or hepatocytes in human and animals and does not enhance of inhibit the growth of HeLa cells. Because of such nature, the growth of vascular endothelial cells can be enhanced selectively and, as the results, new formation of blood vessels can be effected smoothly without causing secondary reactions.

The term "it does not enhance the growth of fibroblasts; vascular smooth muscle cells or hepatocytes and does not enhance or inhibit the growth of HeLa cells" as used herein includes two cases; one case meaning that it does not enhance the growth of fibroblasts, vascular smooth muscle cells or hepatocytes and does not enhance or inhibit the growth of HeLa cells at all, and the other case meaning that it shows these activities to some extent but to an extremely small degree in comparison with its activity to enhance the growth of vascular endothelial cells.

Illustrative procedures for the measurement of activities of the protein of the present invention to enhance the growth of vascular endothelial cells, fibroblasts, vascular smooth muscle cells, hepatocytes and HeLa cells and to inhibit the growth of HeLa cells will be described later in detail in Examples.

In addition to the above properties, the protein of the present invention shows an affinity for concanavalin A. In the present invention, the affinity for concanavalin A was examined in the following manner.

Measurement of affinity for concanavalin A

Using a dot blot apparatus (BioDot; Bio-Rad Laboratories, Inc.), a 500 ng portion of the purified product described in the foregoing was adsorbed to a nitrocellulose membrane (Bio-Rad Laboratories, Inc.) which has in advance been soaked in 10 mM Tris-HCl buffer (pH 7.5) containing 0.15 M NaCl. After air-drying, the resulting membrane was washed by soaking it for 10 minutes in 10 mM Tris-HCl buffer (pH 7.5) containing 0.15 M NaCl and 0.05% Tween and then replacing the washing buffer by a fresh one. After repeating the washing step 4 times, the membrane was soaked for 1 hour at 4°C in the same buffer which has been further supplemented with 1% BSA (bovine serum albumin), and washed again.

The thus treated membrane was soaked in a solution containing 10 μ g/ml of labelled horseradish peroxidase (HRP) - concanavalin A at 4°C for 1 hour and washed again. Thereafter, the HRP remaining after the washing was allowed to perform a coloring reaction in the presence of H_2O_2 using 3,3'-diaminobenzidine as a substrate, in order to judge the affinity of the inventive protein for concanavalin A. As the results, the purified product blotted on the membrane showed development of a brown color, while a control test resulted in no coloration, thus confirming the affinity of the purified product for concanavalin A.

As described in the foregoing, the protein of the present invention is possessed of excellent ability to enhance vascular endothelial cells growth as well as its function to enhance new formation of blood vessels. Because of such nature, a physiologically active pharmaceutical preparation containing the inventive protein can be used as a healing enhancer of wound, burn injury, decubitus, postoperative tissue damage or the like or as a drug for the treatment of cardiac angiopathy, as well as its application to artificial organs such as artificial blood vessel, artificial skin and the like. In addition, antibodies specific for the protein of the present invention and inhibitors of the inventive protein can be used effectively as diagnostic and therapeutic drugs of malignant tumor, retinopathy, chronic rheumatoid arthritis and the like.

€ EXAMPLES

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The following examples are provided to further illustrate the preparation process of the protein of the present invention, the measurement of its molecular weight, its activities on various cells and the presence or absence of its sugar chain moiety. It is to be understood, however, that the examples are for purpose of illustration only and are not intended as a definition of the limits of the invention.

Example 1

(A) Preparation of the protein, measurement of its molecular weight and determination of its aminoacid sequence

(1) To 10 liters of HUOCA-III serum-free culture supernatant was added CHAPS (3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate; Dojin Kagaku K.K.) to a final concentration of 0.03%. The thus prepared serum-free culture supernatant was applied to a 40 ml volume of S-Sepharose (Fast Flow, Pharmacia) which has been equilibrated in advance with 10 mM phosphate buffer (pH 7.2) containing 0.15 M NaCl and 0.03% CHAPS, and the contents were adsorbed at a flow rate of 200 ml/hour at 4°C. After washing with the just described buffer solution containing 0.15 M NaCl, the adsorbed contents were eluted by a linear NaCl gradient using two buffers containing 0.15 M NaCl and 2.0 M NaCl at a flow rate of 200 ml/hour

and at a temperature of 4°C. The eluate was checked for its absorbance at 280 nm and collected as fractions of 6.7 ml/tube. Results of the absorbance measurement at 280 nm are shown in Fig. 1.

Each of the thus collected fractions was checked for its activity to enhance the growth of bovine aorta endothelial cells in the following manner. As shown in Fig. 2, the cell growth enhancing activity was found mostly in fractions 12 to 24.

Measurement of activity to enhance the growth of bovine aorta endothélial cells

Bovine aorta endothelial cells were suspended in DME (Dulbecco's Modified Eagle's) medium (Flow Laboratories, Inc.) which has been supplemented with 10% fetal calf serum, and the cell suspension was poured in a 24 well multi-dish (Corning Glassworks) with a density of 5 x 10³ cells/well. On the following day, the medium was replaced by fresh DME medium containing 5% fetal calf serum, and a sample to be tested was added to the fresh medium, followed by 4 days of culturing to measure the number of resulting cells.

(2) The fractions obtained in the above step (1) having high vascular endothelial cell growth-enhancing activities were pooled and diluted with a buffer solution by a factor of 3, and the contents were adsorbed to heparin-Sepharose CL-6B (Pharmacia; bed volume, 4 ml) which has been equilibrated in advance with a buffer solution containing 0.5 M NaCl, at a flow rate of from 0.2 to 0.4 ml/minute and at a temperature of 4°C. After washing with the same buffer solution containing 0.5 M NaCl, the adsorbed contents were eluted by a linear NaCl gradient using two buffers containing 0.5 M NaCl and 2.0 M NaCl at a flow rate of 0.2 ml/min and at a temperature of 4°C. The eluate was checked for its absorbance at 280 nm and collected as fractions of 3 ml/tube. Results of the absorbance measurement at 280 nm are shown in Fig. 3.

Each of the thus collected fractions was checked for its activity to enhance the growth of bovine aorta endothelial cells in the same manner as described above. As shown in fig. 4, the cell growth enhancing activity was found mostly in fractions 23 to 30.

(3) The fractions obtained in the above step (2) having high vascular endothelial cell growth-enhancing activities were pooled and diluted with a buffer solution by a factor of 3, and the contents were adsorbed on to a TSK-heparin 5PW column (7.5 mm in inside diameter and 7.5 cm in length; Tosoh Corp.) which has been equilibrated in advance with a buffer solution containing 0.5 M NaCl. After washing with the same buffer solution containing 0.5 M NaCl, the adsorbed contents were eluted by a linear NaCl gradient using two buffers containing 0.5 M NaCl and 2.0 M NaCl, at a flow rate of 0.5 ml/min and at room temperature. The eluate was checked for its absorbance at 215 nm and collected as fractions of 0.5 ml/tube. Results of the absorbance measurement at 215 nm are shown in Fig. 5.

Each of the thus collected tractions was checked for its activity to enhance the growth of bovine aorta endothelial cells in the same manner as described above. As shown in Fig. 6, the cell growth enhancing activity was found mostly in fractions 30 to 32.

(4) The fractions obtained in the above step (3) having high vascular endothelial cell growth-enhancing activities were pooled and subjected to reverse phase chromatography using a vp-318 column (4.6 mm in inside diameter and 30 mm in length; Senshu Kagaku Co., Ltd.). In the presence of 0.1% trifluoroacetic acid (TFA), a linear gradient elution was carried out by increasing the concentration of acetonitrile from 10% to 60%, at a flow rate of 1.0 ml/min. The eluate was checked for its absorbance at 215 nm and collected as fractions of 10 ml/tube. Results of the absorbance measurement at 215 nm are shown in Fig. 7.

Each of the thus collected fractions was checked for its activity to enhance the growth of bovine aorta endothelial cells in the same manner as described above, with the results shown in Fig. 8. By collecting peak fractions, a highly purified product having high vascular endothelial cell growth-enhancing activity was obtained.

(5) The molecular weight of the highly purified product obtained in the above step (4) was measured by SDS polyacrylamide gel electrophoresis.

The following 6 authentic samples whose molecular weights have been confirmed were used as molecular weight markers, and the electrophoresis was carried out in the same manner as described in the foregoing.

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[Molecular weight markers]								
Rabbit muscle phosphorylase	(M.W., 97,400 daltons)							
2. Bovine serum albumin	(M.W., 66,200 daltons)							
3. Ovalbumin	(M.W., 45,000 daltons)							
4. Carbonic anhydrase	(M.W., 31,000 daltons)							
5. Soybean trypsin inhibitor	(M.W., 21,500 daltons)							
6. Lysozyme	(M.W., 14,400 daltons)							

The thus obtained electrophoresis pattern is shown in Fig. 9. As is evident from the figure, the highly purified product obtained in the above step (4) has a molecular weight of 72,000 to 80,000 daltons under non-reducing condition, or 79,000 to 85,000 daltons under reducing condition, when measured by SDS polyacrylamide gel electrophoresis. It is evident also that the purified product is a single chain protein.

After the electrophoresis, the gel was cut out at intervals of 2 mm. Each of the thus cut portions was put into a test tube, ground into pieces, mixed with $500 \,\mu$ l of a buffer solution 0.03% CHAPS, 20 mmol PB pH 7.2 and then shaken at 4°C for 16 hours. The resulting mixture was centrifuged to recover supernatant fluid which was subsequently dialyzed against a buffer solution 0.03% CHAPS, 20 mmol PB pH 7.2. Contents in the thus dialyzed solution was freeze-dried and then dissolved in 100 μ l of a buffer solution 0.03% CHAPS, 20 mmol PB pH 7.2 to measure the activity to enhance the growth of bovine aorta endothelial cells in the same manner as described in the foregoing. As shown in Figure 10, the endothelial cell growth-enhancing activity was observed in 72,000-80,000 molecular weight fraction obtained under non-reducing condition.

When the amino acid sequence of the highly purified product was determined in accordance with the procedure described in the foregoing, it was confirmed that the product contained three peptide chains respectively represented by the Sequence ID Nos. 1, 2 and 3.

Also, in order to confirm the addition of sugar chains to the highly purified product, 5 μ l (250 ng) of the high purity product and 3.2 μ l of N-glycanase (Genzyme Corp.; 250 units/ml) were added to 30 μ l of 50 mM Tris-HCl buffer (pH 8.0). After 18 hours of reaction, the resulting mixture was subjected to 0.1% SDS-10% polyacrylamide gel electrophoresis, followed by silver staining. As shown in Fig. 11, the resulting electrophoresis pattern clearly indicated a decrease in the molecular weight of the N-glycanase-treated product due to the separation of sugar chains.

(B) Cloning of the DNA and estimation of the amino acid sequence

(a) Synthesis of the cCNA

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A 5 μ l portion of the total RNA sample (10 μ g/ μ l) which has been prepared from the human ovarian tumor cell line HUOCA-III by the SDS-phenol method was incubated at 70°C for 5 minutes and then cooled down rapidly. After 5 minutes of cooling on an ice bath, to this were added 10 μ l of a 5 x buffer solution for reverse transcription use (250 mM Tris-HCl/pH 8.3, 375 mM KCl, 15 mM MgCl2), 15 μ l of 2.5 mM dNTP (a mixture of dATP, dCTP, dGTP and dTTP; Takara Shuzo Co., Ltd.), 0.5 μ l of 1 M DTT (dithiothreitol), 1 μ l of oligo(dT)₁₂₋₁₈ (Amersham), 2.5 μ l of a ribonuclease inhibitor (200 U/ μ l, Takara Shuzo Co., Ltd.), 13 μ l of distilled water and 3 μ l of M-MLV reverse transcriptase (200 U/ μ l, GIBCO-BRL). The thus prepared mixture was incubated at 37°C for 1 hour to effect cDNA synthesis. After removing the proteinous materials from the resulting reaction mixture by phenol treatment, the cDNA of interest was recovered by ethanol precipitation, dissolved in 50 μ l of distilled water and then stored at -80°C.

(b) Amplification of the cDNA which encodes the HUOCA-III-originated novel protein by polymerase chain reaction (PCR)

To 5 μ l of the cDNA aqueous solution were added 70 μ l of distilled water, 10 μ l of a 10 x buffer solution for PCR use (500 mM KCl, 15 mM MgCl2, 100 mM Tris-HCl/pH 8.3, 0.01% (w/v) gelatin), 8 μ l of dNTP (Takara Shuzo Co., Ltd.), 3 μ l of a 5' primer (5' TCTTTTAGGCACTGACTCCGAACAGGATTCTTTCAC 3', 1 μ g/ μ l) and 3 μ l of a 3' primer (5' GTTGTATTGGTGGATCCTTCAGACACACTTACTTCAG 3'). The thus prepared mixture was incubated at 95°C for 7 minutes, followed by rapid cooling. The thus treated solution was mixed with 1 μ l

of Ampli Taq DNA polymerase (5 U/µl, Perkin Elmer Cetus), and the surface of the reaction solution was covered with mineral oil (nujol mineral oil manufactured by Perkin Elmer Cetus). Thereafter, PCR was carried out by 30 repetitions of a three step reaction (94°C for 1 minute, 60°C for 2 minutes and 72°C for 3 minutes). After completion of the reaction, mineral oil was removed by chloroform treatment, proteinous materials were removed by phenol treatment and then the PCR product was recovered by ethanol precipitation.

(c) Digestion of the PCR product with BamHI

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An 85 μ l portion of the PCR product was mixed with 10 μ l of a 10 x buffer solution for BamHI reaction use (1.5 M NaCl, 60 mM Tris-HCl/pH 7.9, 60 mM MgCl2) and 5 μ l of an aqueous solution of BamHI (15 U/ μ l, Nippon Gene), and the resulting mixture was incubated at 37°C for 1 hour.

(d) Purification of the BamHI-digested PCR product

The PCR product thus digested with BamHI was subjected to 0.7% agarose gel electrophoresis at a constant voltage (100 V). After completion of the electrophoresis, the gel was stained with ethidium bromide to observe DNA bands using a UV transilluminator. A portion of the gel where a DNA band of 2.3 kb was observed was cut out, and the PCR product in the cut portion was purified using Sephaglas Band Prep Kit (Pharmacia).

(e) Digestion of the pUC18 plasmid vector with BamHI

A 2 μ l portion of pUC18 solution (1 μ g/ μ l, Takara Shuzo Co., Ltd.) was mixed with 6.6 μ l of distilled water, 3 μ l of the 10 x buffer solution for BamHI reaction use and 1.4 μ l of BamHI (15 U/ μ l, Nippon Gene), and the resulting mixture was incubated at 37°C for 1 hour to digest the plasmid. After completion of the reaction, proteinous materials were removed by phenol treatment and the thus digested plasmid fragments were recovered by ethanol precipitation. The thus recovered plasmid fragments were dissolved in 33 μ l of distilled water and mixed with 4 μ l of CIP buffer (50 mM Tris-HCl/pH 8.0, 1 mM MgCl₂) and 3 μ l of alkaline phosphatase (calf intestine origin, 2,500 U/ml, Toyobo Co., Ltd.). The resulting mixture was incubated at 37°C for 40 minutes and then at 50°C for 20 minutes. After completion of the reaction, the BamHI-digested fragments of the plasmid vector pUC18 were recovered by phenol treatment and subsequent ethanol treatment.

(f) Transformation of E.Coli JM109 with the PCR product

To 6 μ l (30 μ g) of the the BamHl-digested PCR product were added 2 μ l (200 μ g) of the pUC18 digest prepared in the above step (e), 2 μ l of a 10 x ligation buffer solution (10 mM ATP, 200 mM DTT, 100 mM MgCl₂, 500 mM Tris-HCl/pH 7.9), 9 μ l of distilled water and 1 μ l of T4 DNA ligase (500 U/ μ l, Nippon Gene). After overnight reaction at 16°C, a portion of the resulting reaction solution was added to 100 μ l of a suspension of E. Coli JM109 competent cells (Nippon Gene). The thus prepared mixture was allowed to stand still for 20 minutes on an ice bath, heat-treated at 42°C for 45 seconds and then allowed again to stand still on an ice bath for at least 2 minutes. The thus treated mixture was added to 400 μ l of High-compitence broth (Nippon Gene) and stirred on a shaker at 37°C for 60 minutes. To this were added 40 μ l of 2% X-Gal (5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside) dissolved in diethylformamide and 40 μ l of 100 mM IPTG (isopropyl- β -D-thio-galactopyranoside). The thus prepared mixture was poured on LB plate medium (0.5% yeast extract, 1% Bacto-Trypton, 1.5% agar, 1% NaCl, 50 μ g/ml ampicillin, pH 7.5) and incubated overnight at 37°C to find white (recombinant) colonies and blue (non-recombinant) colonies grown on the medium. By isolating white colonies, a JM109 transformant into which the cDNA of interest has been inserted was selected.

(g) Preparation of the plasmid

The plasmid-introduced JM109 was cultured overnight at 37°C in 100 ml of LB medium (1% Bacto-Trypton, 0.5% yeast extract, 1% NaCl, pH 7.5). When the cells reached their logarithmic growth phase, they were collected by centrifugation (5 minutes, 5,000 pm, 0°C) and suspended in 4 ml of P1 buffer solution (100 µg/ml RNase A, 50 mM Tris+HCl/pH 8.0, 10 mM EDTA). The resulting cell suspension was mixed with 4 ml of P2 buffer solution (200 mM NaOH, 1% SDS) to carry out an alkali treatment at room temperature for 5 minutes. After the alkali denaturation, the resulting mixture was neutralized by adding 4 ml of P3 buffer solution (2.55 mM Potassium acetate, pH 4.8) and then centrifuged at 15,000 rpm for 30 minutes at 4°C. The thus obtained supernatant fluid was applied to a QIAGEN-MIDI column-pack 100 (DIAGEN) which has been equilibrated in advance with 2 ml of QB buffer solution (750 mM NaCl, 50 mM MOPS [3-(N-morpholino)propanesulfonic acid]/pH

7.0, 15% ethanol). After washing the column twice with 4 ml of QC buffer solution (1 M NaCl, 50 mM MOPS/pH 7.0, 15% ethanol), the plasmid was eluted with 2 ml of QF buffer solution (1.2 M NaCl, 15% ethanol, 50 mM MOPS/pH 8.0). The eluate was mixed with 500 μ l of isopropanol and centrifuged at room temperature for 30 minutes. Thereafter, the precipitate thus obtained was washed with 70% ethanol and dissolved in 100 μ l of distilled water.

(h) Determination of the nucleotide sequence by the dideoxy method

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A 16 µl (3 µg) portion of the plasmid solution prepared in the above step (g) was mixed with 2 µl of 2 N NaOH and 2 μl of 2 mM EDTA, and the mixture was incubated at 37°C for 25 minutes to denature the plasmid. After the alkali denaturation, the resulting solution was mixed with 2 µl of 3 M sodium acetate and 100 µl of cold ethanol, and ethanol precipitation was effected by maintaining the mixture for 10 minutes at -80°C. The thus precipitated plasmid was recovered by centrifugation, washed with 70% ethanol and then dissolved in 7 μl of distilled water. To this were added 1 μl of a primer (0.5 pmole) and 2 μl of a 5 x buffer solution A (250 mM NaCl, 200 mM Tris-HCl/pH 7.5, 100 mM MgCl2). After 2 minutes of incubation at 65°C, the resulting solution was gradually cooled down to 30°C to effect annealing of the denatured plasmid and the primer. To the resulting solution were added 1 μl of 0.1 M dithiothreitol, 2 μl of a labeling mixture (1.5 μM 7-deaza-dGTP, 1.5 μM dATP, 1.5 μM dTTP), 0.5 μl of [α-35S]dCTP (1,000 Ci/mmole, Amersham) and 2 μl of Sequenase Ver. 2.0 (1.5 U/μl, United States Biochemical Corporation). After 5 minutes of reaction at 37°C, a 3.5 µl portion of the resulting reaction mixture was added to 2.5 μl of each of a G solution (80 μM 7-deaza-dGTP, 80 μM dATP, 80 μM dCTP, 80 μM dTTP, 8 μM ddGTP, 50 mM NaCl), an A solution (80 μM 7-deaza-dGTP, 80 μM dATP, 80 μM dCTP, 80 μΜ dTTP, 8 μΜ ddATP, 50 mM NaCl), a C solution (80 μΜ 7-deaza-dGTP, 80 μΜ dATP, 80 μΜ dCTP, 80 μΜ dTTP, 8 μM ddCTP, 50 mM NaCl) and a T solution (80 μM 7-deaza-dGTP, 80 μM dATP, 80 μM dCTP, 80 μM dTTP, 8 μM ddTTP, 50 mM NaCl). In this instance, each of these solutions was kept at 37°C prior to its use. After 5 minutes of reaction at 37°C, the reaction was terminated by adding 4 µl of a reaction termination solution (95% formamide, 0.05% Bromophenol Blue, 20 mM EDTA, 0.05% Xylene Cyanol FF). Thereafter, the reaction mixture was heated at 90°C for 5 minutes, followed by rapid cooling, and a 2.5 µl portion of the resulting sample was subjected to electrophoresis. In this case, a composition consisting of 7 M urea, 10% HydroLink™ LONG-RANGER (AT Biochem), 100 mM Tris-HCl, 100 mM boreic acid and 2 mM EDTA was made into gel using 0.05% ammonium persulfate and 0.0005% N,N,N',N'-tetramethylenediamine (TEMED), and the electrophoresis was carried out at a constant power of 60 W using a TEB buffer (50 mM Tris, 50 mM boreic acid, 1 mM EDTA). After completion of the electrophoresis, the gel was dried on a filter paper and subjected to autoradiography to determine the nucleotide sequence of the DNA of interest.

The thus determined DNA sequence is shown in the Sequence ID No. 5, and an amino acid sequence deduced from the DNA sequence is shown in the sequence ID No. 4.

As generally known in this art, the amino acid sequence shown in the Sequence ID No. 4 has a signal peptide. Therefore, the protein of the present invention may be the whole Sequence ID No. 4, a portion of the sequence (for example, the Sequence ID No. 4 except the sequence of a signal peptide), or the portion of the Sequence together with a linker.

The protein of the present invention includes at least an active portion having an activity to enhance the growth of vascular endothelial cells obtainable from a nucleotide sequence or a portion of the nucleotide sequence represented by the Sequence ID No. 5. The DNA corresponding to the signal peptide in the nucleotide sequence represented by the Sequence ID No. 5 may be changed another DNA corresponding to another signal peptide, if necessary, a signal peptide together with a linker DNA sequence may be used in the DNA fragment represented by the Sequence ID No. 5 attached hereto.

Example 2 Affinity for concanavalin A

The highly purified product obtained in the step (4) of Example 1 was checked for its affinity for concanavalin A in accordance with the procedure described in the foregoing. As the results, it was confirmed that the purified product was possessed of the affinity for concanavalin A, which is a

In addition, on the basis of the results obtained in Examples 1 and 2, it was confirmed that the high purity product of the step (4) was a single chain glycoprotein.

Example 3 New formation of blood vessels

A total of 10 avian eggs, fertilized for 8 days, were used in each test group. A filter (6 mm in diameter) which has been impregnated with a varied amount of the highly purified product (glycoprotein of this invention) ob-

tained in the step (4) of Example 1 was put on the chorio-allantoic membrane of each egg. After 3 days of incubation at 37°C under a moist condition, new formation of blood vessels was observed under a stereoscopic microscope. The judgement was made as positivre (+, new formation of blood vessels around the filter) or negative (-, no formation of new blood vessels), and the number of positive eggs in each test group was counted. As a comparative example, the same experiment was carried out except that the filter was impregnated with physiological saline instead of the purified product. The results are shown in Table 1.

Table 1

Test group	Amount of glycoprotein	Positive effs/Total
1 1	0 (physiological saline)	0/10
2	1 ng/filter	1/10
3	10 ng/filter	3/10
4	50 ng/filter	5/10
5	100 ng/filter	6/10

It is evident from the above table that the glycoprotein of the present invention is possessed of a function to enhance new formation of blood vessels.

Example 4 Growth enhancing effect on human umbilical cord vascular endothelial cells

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Human umbilical cord vascular endothelial cells were prepared in the usual way and inoculated into a collagen-coated 24 well multi-dish (Corning Glassworks) with a cell density of 1 x 10⁴ cells/well, using MCDB107 medium (Kyokuto Pharmaceutical Industrial Co., Ltd.) supplemented with 20% fetal calf serum. At intervals of 2 days from the next day, the medium was exchanged for a fresh medium containing 5% fetal calf serum and a predetermined amount (see Table 2) of the glycoprotein of the present invention obtained in the step (4) of Example 1. The number of cells was counted on the eighth day, with the results shown in Table 2.

Table 2

Table 2									
Glycoprotein (ng/ml)	Cell count (cells/well)								
0	27168								
0.3	29460								
1.0	30920								
3.3	37492								
10.0	43072								
33.3	54772								
100.0	53988								
333	46460								

As is evident from the above table, the glycoprotein of the present invention is possessed of a function to enhance the growth of human umbilical cord vascular endothelial cells.

Example 5 Presence/absence examination of growth enhancing effect on fibroblasts

A primary culture of human dermis fibroblasts prepared from human skin was subcultured, and the eighth subculture was inoculated into a 24 well multi-dish with a cell density of 5 x 10³ cells/well, using DME medium (Flow Laboratories, Inc.) supplemented with 10% fetal calf serum. At intervals of 2 days from the next day, the medium was exchanged for fresh DME medium containing 0.5% fetal calf serum and 100 ng/ml of the glycoprotein of the present invention obtained in the step (4) of Example 1.

As a comparative example, the same procedure was repeated except that the glycoprotein was eliminated

from the medium or a basic fibroblast growth factor (bFGF) was used in an amount of 1 ng/ml instead of the glycoprotein.

The number of cells was counted on the eighth day, with the results shown in Table 3.

Table 3

Component added	Cell count on 8th day (cells/well)
No addition	28248
Glycoprotein of Example 1	24325
bFGF	42645

As is evident from the above table, bFGF strongly enhances the growth of fibroblasts, but the number of fibroblasts on the eighth day in the case of the addition of the glycoprotein of the present invention obtained in Example 1 is almost the same as that of the case of the control (no addition), thus showing that the inventive glycoprotein hardly has a function to enhance the growth of fibroblasts.

Example 6 Presence/absence examination of growth enhancing effect on vascular smooth muscle cells

A primary culture of human smooth muscle cells prepared from an umbilical cord was subcultured, and the sixth subculture was inoculated into a 24 well multi-dish with a cell density of 5 x 10³ cells/well, using DME medium supplemented with 10% fetal calf serum. At intervals of 2 days from the next day, the medium was exchanged for fresh medium containing 100 ng/ml of the glycoprotein of the present invention obtained in the step (4) of Example 1.

As a comparative example, the same procedure was repeated except that the glycoprotein was eliminated from the medium or a basic fibroblast growth factor (bFGF) was used in an amount of 1 ng/ml instead of the glycoprotein.

The number of cells was counted on the eighth day, with the results shown in Table 4.

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Table 4

Component added	Cell count on 8th day (cells/well)
No addition	6192
Glycoprotein of Example 1	7480
bFGF	48962

As is evident from the above table, the number of smooth muscle cells on the eighth day in the case of the addition of the glycoprotein of the present invention obtained in Example 1 is almost the same as that of the case of the control (no addition), thus showing that the inventive glycoprotein has no activity to enhance the growth of human smooth muscle cells.

Example 7 Presence/absence examination of growth enhancing effect on hepatocytes

Hepatic parenchymal cells (to be referred to as "hepatocytes" hereinafter) were prepared in accordance with the procedure of Takahashi et al. (Tissue Culture, vol.12, No.8, pp.308 - 312, 1986). The thus prepared hepatocytes were suspended in an inoculation medium (WE basal medium supplemented with 5% fetal calf serum and 10-8 M dexamethasone) to a cell density of 5.0 x 104 cells/0.2 ml, and the resulting hepatocyte suspension was inoculated into a collagen-coated 24 well multi-dish. After 4 hours of the culturing, the medium was replaced by WE basal medium and the glycoprotein of the present invention obtained in Example 1 was added to the fresh medium in a predetermined amount as shown in Table 5. The same process was repeated after additional 16 hours of the culturing. The medium was exchanged again for fresh WE basal medium 40 hours after the commencement of the culturing, and 3H-thymidine was added to the fresh medium to carry out 2 hours of pulse-labeling. After completion of the pulse-labeling, the culture supernatant was removed, and the remaining cells were washed with a cold phosphate buffer (PBS), 2% perchlorate and 95% cold ethanol in that order and then dried at room temperature. In this instance, each washing step was repeated three times. The thus dried cells in each well were lysed by adding 0.8 ml of a 1% SDS/0.1 N NaOH solution and maintaining

the mixture at 37°C for at least 1 hour. A 0.5 ml portion of the resulting lysate was pipetted off from each well and put into a scintillation vial. Thereafter, the content in the vial was mixed with 7 ml of a scintillator (OptiFlow, Packard), and the radioactivity was measured using a scintillation counter to examine ³H-thymidine uptake.

As a comparative example, the same experiment was carried out except that a mixture of insulin (100 nM/ml) and epidermal growth factor (EGF, 50 ng/ml) was used instead of the glycoprotein of the present invention.

The results are shown in Table 5.

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Table 5

Component added	Uptake of ³ H-thymide					
Glycoprotein of Example 1						
300 ng/ml	5697 DPM					
100 ng/ml	4347 DPM					
30 ng/ml	4869 DPM					
10 ng/ml	4619 DPM					
Insulin + EGF	76815 DPM					
(100 nM + 50 ng/ml)						
Control (no addition)	4992 DPM					

As is evident from the above table, uptake of ³H-thymidine does not occur by the addition of the glycoprotein of the present invention, thus showing that the inventive glycoprotein has no activity to enhance the growth of hepatocytes.

Example 8 Presence/absence examination of growth enhancing or inhibiting effect on HeLa cells

HeLa-S3 cells were suspended in MEM medium containing 5% bovine serum to a cell density of 1 x 10^5 cells/ml. The thus prepared HeLa-S3 cell suspension was dispensed in $100~\mu$ portions into wells of a 96 well multi-dish. After 24 hours of culturing, the resulting medium was replaced by fresh MEM medium which has been supplemented writh 5% fetal calf serum and a predetermined amount of the glycoprotein obtained in Example 1, and the culturing was continued for additional 48 hours.

Since the presence or absence of the growth inhibiting effect was not able to be judged clearly with the naked eye under a phase-contrast microscope, the judgement was made by staining the cells with Crystal Violet. That is, each well of the dish after the culturing was washed with a phosphate buffer and then filled with a 10% formalin solution for a period of 30 minutes to fix the cells. The thus treated dish was dried after washing it with running water to remove formalin, and the cells in the dish were stained for 15 minutes with a 0.2% Crystal Violet solution containing 2% ethanol. After removing unbound pigment by washing the dish in running water, and subsequently drying the dish, a predetermined amount of 1% sodium dodecyl sulfate solution was added to each well to dissolve the bound pigment. Thereafter, absorbance of the thus dissolved Crystal Violet was measured at a wave length of 540 nm.

As a control, the same culturing step was repeated except that the glycoprotein was not used, and the Crystal Violet staining and absorbance measurement at 540 nm were carried out in the same manner.

The results are shown in Table 6 in which the absorbance of the control at 540 nm is expressed as 1.00.

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Table 6

Component added	Ratio of absorbance at 540 nm					
Glycoprotein of Example 1						
300 ng/ml	1.02					
100 ng/ml	1.01					
30 ng/ml	1.01					
10 ng/ml	1.02					
Control (no addition)	1.00					

As shown in the above table, the absorbance at 540 nm hardly changed by the addition of the glycoprotein of the present invention in comparison with the case of the control (no addition), thus confirming that the inventive glycoprotein has no activity to enhance or inhibit the growth of HeLa cells.

Example 9 Migration-stimulating activity on vascular endothelial cells and smooth muscle cells

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Primary culturing of vascular endothelial cells was carried out by isolating the cells from rabbit cornea capillary vessels in the usual way. The migration-stimulating activity of the cells was measured in accordance with the Boyden's test using Boyden's chamber. That is, DME medium supplemented with 10% fetal calf serum and a predetermined amount of the glycoprotein obtained in Example 1 was put into the lower compartment of the Boyden's chamber, and another DME medium supplemented with 10% fetal calf serum and 2 x 104/ml of vascular endothelial cells was put into the upper compartment of the chamber. Thereafter, culturing was carried out at 37°C for 4 hours.

A similar test was carried out using primary-cultured smooth muscle cells which have been isolated from rat pulmonary artery

After the culturing, the thus treated cells were stained with Diff-Quick solution, and the number of migrated cells per visual field was counted under a microscope, with the results shown in Table 7.

Table 7

lable /									
	The number of migrated cells								
Glycoprotein	Vascular endothelial cells	Smooth muscle cells							
300 ng/ml	268	0							
100 ng/ml	50	0							
30 ng/ml	37	0							

As is evident from the above table, the glycoprotein of the present invention shows migration-stimulating activity on vascular endothelial cells but not on smooth muscle cells.

Thus, it is apparent that there has been provided, in accordance with the present invention, a novel protein of human origin, as well as a process for the production thereof. Since the protein of the present invention enhances the growth of vascular endothelial cells but does not activate the growth of smooth muscle cells, fibroblasts and hepatocytes and also does not enhance or inhibit the growth of HeLa cells, it can enhance the growth of vascular endothelial cells selectively and therefore can enhance new formation of blood vessels smoothly without causing secondary reactions. Because of such excellent properties, especially its activity to enhance new formation of blood vessels, the protein of the present invention can be applied to a healing enhancer of wound, burn injury, decubitus, postoperative tissue damage or the like or as a drug for the treatment of cardiac angiopathy, as well as its application to artificial organs such as artificial blood vessel, artificial skin and the like. It also can be applied to diagnostic and therapeutic drugs of malignant tumor, retinopathy, chronic rheumatoid arthritis and the like.

In addition, the protein of the present invention can be obtained with a high productivity and a high purity in comparison with the prior art physiologically active factors.

SEQUENCE LISTING

5	(1) GENERAL INFORMATION:
10	 (i) APPLICANT: (A) NAME: TERUMO KABUSHIKI KAISHA (B) STREET: 44-1, Hatagaya 2-chome, Shibuya-ku (C) CITY: TOKYO (E) COUNTRY: JAPAN (F) POSTAL CODE (ZIP): 151
15	
	(ii) TITLE OF INVENTION: Novel protein of human origin and its production process
20	(iii) NUMBER OF SEQUENCES: 7(iv) COMPUTER READABLE FORM:(A) MEDIUM TYPE: Floppy disk
25	(B) COMPUTER: IBM PC compatible(C) OPERATING SYSTEM: PC-DOS/MS-DOS(D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
30	(v) CURRENT APPLICATION DATA: APPLICATION NUMBER: EP 92 403 199.0 (vi) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 3-337999
35	(B) FILING DATE: 28-NOV-1991
	(2) INFORMATION FOR SEQ ID NO: 1:
40	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 7 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: peptide
	(iii) HYPOTHETICAL: NO
50	(v) FRAGMENT TYPE: N-terminal
55	<pre>(vi) ORICINAL SOURCE: (A) ORGANISM: Homo sapiens (G) CELL TYPE: Ovarian (H) CELL LINE: HUOCA II / HUOCA III</pre>

	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 1:
5	Arg 1	Asn Thr Ile His Glu Phe 5
	(2) INFO	RMATION FOR SEQ ID NO: 2:
10	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 10 amino acids (B) TYPE: amino acid
15		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: peptide
20	(iii)	HYPOTHETICAL: NO
	(v)	FRAGMENT TYPE: internal
25	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Homo sapiens (G) CELL TYPE: Ovarian (H) CELL LINE: HUOCA II / HUOCA III
30	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 2:
35	Glu 1	Phe Gly His Glu Phe Asp Leu Tyr Glu 5 10
، د مدهنجین	(2) INFO	RMATION FOR SEQ ID NO: 3:
40	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear
45	(ii)	MOLECULE TYPE: peptide
	(iii)	HYPOTHETICAL: NO
50	(v)	FRAGMENT TYPE: C-terminal
5 5	(vi)	QRIGINAL SOURCE: (A) ORGANISM: Homo sapiens (G) CELL TYPE: Ovarian (H) CELL LINE: HUOCA II / HUOCA III

	(ix) FEATURE:
	(A) NAME/KEY: Modified-site
5	(B) LOCATION: 3
	(D) OTHER INFORMATION: /label= Xaa
	/note= "unidentified amino acid residue"
	dela lesidue
10	(ix) FEATURE:
	(A) NAME/KEY: Modified-site
	(B) LOCATION: 10
	(D) OTHER INFORMATION: /label= Xaa
15	<pre>/note= "unidentified amino acid residue"</pre>
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
20	Glu Ser Xaa Val Leu Thr Ala Arg Gln Xaa Phe Pro Ser Arg Asp Leu
	1 5 10 15
	(2) INFORMATION FOR SEQ ID NO: 4:
25	TON DEG ID NO; 4;
	(1) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 728 amino acids
	(B) TYPE: amino acid
30	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: protein
35	(iii) HYPOTHETICAL: YES
	(wi) Ontorwo
	(vi) ORIGINAL SOURCE:
	(A) ORGANISM: Homo sapiens
40	(G) CELL TYPE: ovarian
	(H) CELL LINE: HUOCA II / HUOCA III
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
45	The second of th
	Met Trp Val The Los Law Law D
	Met Trp Val Thr Lys Leu Leu Pro Ala Leu Leu Cln His Val Leu
	1 5 10 15
50	Leu His Leu Leu Leu Pro Ile Ala Ile Pro Tyr Ala Glu Gly Gln
	20 25 30
	Arg Lys Arg Arg Asn Thr Ile His Glu Phe Lys Lys Ser Ala Lys Thr
	35 40 45
55	

	me	50	TTE	Lys	116	Asp	Pro 55	Ala	Leu	Lys	Ile	Lys 60	Thr	Lys	Lys	Val
5	Asn	Thr	Ala	Asp	Gln	Cys		Asn	Arg	Cys	Thr		Asn	Lvs	Glv	Len
	65					70	,		J	·	75			~J C		80
	Pro	Phe	Thr	Cys	Lys	Ala	Phe	Val	Phe	Asp	Lys	Ala	Arg	Lys	Gln	Cys
10					85					90					95	
	Leu	Trp	Phe	Pro	Phe	Asn	Ser	Met	Ser	Ser	Gly	Val	Lys	Lys	Glu	Phe
				100					105					110		
	Gly	His		Phe	Asp	Leu	Tyr	Glu	Asn	Lys	Asp	Tyr	Ile	Arg	Asn	Cys
15			115					120					125			
	lle		Gly	Lys	Gly	Arg		Туг	Lys	Gly	Thr	Val	Ser	Ile	Thr	Lys
		130	~-		_		135			•		140				
20		GLY	TTE	Lys	Cys		Pro	Trp	Ser	Ser		Ile	Pro	His	Glu	
	145	Dh.	T	D	c .	150				_	155	_				160
	ser	rne	Leu	Pro		Ser	Tyr	Arg	GLY	Lys	Asp	Leu	Gln	Glu		Tyr
25	Cve	Ana	Acn	Dno	165	C1	C1	01	01. -	170	ρ		•	5 1	175	_
	OJ S	8	กอม	180	urg	ury	ara	ara	185	Gly	Pro	irp	Cys		Thr	Ser
	Asn	Pro	Glu		Arø	Tur	Glu	Vol	-	Asp	Tlo	Dno	Cl-	190	Com.	C1
30			195	•		-3-	GI.	200	Cys	nsp	116	110	205	Cys	Ser	GIU
30	Val	Glu		Met	Thr	Cys	Asn		Glu	Ser	Tvr	Arg	-	Leu	Met	Asn
		210	-			•	215				-3-	220	3			··op
	His	Thr	Glu	Ser	Gly	Lys	_	Cys	Gln	Arg	Trp		His		Thr	Pro
35	225					230					235	·				240
	His	Arg	His	Lys	Phe	Leu	Pro	Glu	Arg	Tyr	Pro	Asp	Lys	Gly	Phe	Asp
					245					250					255	
40	Asp	Asn	Tyr	Cys	Arg	Asn	Pro	Asp	Gly	Gln	Pro	Arg	Pro	Trp	Cys	Tyr
				260					265					270		
	Thr	Leu	Asp	Pro	His	Thr	Arg	Trp	Glu	Tyr	Cys	Ala	Ile	Lys	Thr	Cys
45			275					280					285			
	Ala	Asp	Asn	Thr	Met	Asn	Asp	Thr	Asp	Val	Pro	Leu	Glu	Thr	Thr	Glu
		290					295					300				
	Cys	Ile	Gln	Gly	Gln	Gly	Glu	Gly	Tyr	Arg	Gly	Thr	Val	Asn	Thr	Ile
50	305					310					315					320
	Trp	Asn	Gly	Ile	Pro	Cys	Gln	Arg	Trp	Asp	Ser	Gln	Tyr	Pro	His	Glu
			•		325					330					335	
55	His	Asp	Met	_	Pro	Glu	Asn	Phe	Lys	Cys	Lys	Asp	Leu	Arg	Glu	Asn
				340					345					350		

	Ту	r Cy	s A	rg As	sn P	ro As	sp G	ly S	er G	lu S	er Pi	ro Ti	ים כי	o Pi	ים דו	nr Thr
			35	55				3	60				36		16 11	it inr
5	As	p Pr	o As	sn I	le Aı	g Va	al G			vs Se	er Gl	n II	بر D+	,, ,, ,,		s Asp
		37	0				37			,	- 01	38		U AS	ii C	's Asp
	Me	t Se	r Hi	s Gl	y G1	л As			/r A1	~e~ G1	v Ac	יטע מי		- A-	- m	r Met
10	38	5				39	0	,		. 5 0.			у гу	S AS	n Ty	
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					40	5				41		r cy	s se	r we		
	Lys	s Ası	n Me	t Gl		-	u Hi	c Ar	or Hi			. T	. 01		41	5 p Ala
15				42	0				42		e Pn	e ir	p GI			p Ala
	Sei	Lys	s Le			ıı Acı	n ጥህ	n ()			- 5			43	0	
		•	43	- ··-· 5		u 113	ıı ıy	44:		g AS	n Pro	o Ası			p Al	a His
20	Gly	/ Pro			s Tv	r ፔክ	n (31)				~.	_	449	5		
20	-	450)		- - J		459		n Pr	o re	n II.			o Ası	o Ty	r Cys
	Pro	_		r Ara	z Cve	2 (3)			- m.		_	460				
	465	i			5 934	470	1 013	y AS	p in	r in			· Ile	Val	l Ası	ı Leu
25	Asp	His	Pro	. Val	Tle			. 41.	n I	- 171	475		_			480
	•				485	; ;	. Cys	, WIS	ı Ly:			Gln	Leu	Arg		Val
	Asn	Glv	Ile	Pro	-		. m	۸	. ті.	490					495	5
30		·		500)	VLE	, 1111	· ASI			Trp	Met	Val	Ser	Leu	Arg
00	Tyr	Arg	Asn	-		Tla	Cva	0 1-	505		_			510		
			515	,,	, ,,,,,	116	Cys			, Ser	· Leu	Ile			Ser	Trp
	Val	Leu			Ara	Gla	Carm	520 Dha		•			525			
35		530			••••	GIII			Pro) Ser	Arg		Leu	Lys	Asp	Tyr
			Tro	Leu	ดาง	TIA	535		V-1	***	01	540				
	545			200	ary	550	1112	nsp	vai	His		Arg	Gly	Asp	Glu	Lys
40		Lvs	Gln	Vel	I ou		17-1	.	01		555					560
	.	_, _		vai	565	VSII	vai	ser	GIN	Leu	Val	Tyr	Gly	Pro	Glu	Gly
	Ser	Asn	Lan	Vel		Wa.	1		4.7	570	_				575	
			Deu	580	Leu	met	Lys	Leu		Arg	Pro	Ala	Val	Leu	Asp	Asp
45	Phe	Val	Son	-	T1 -	A			585					590		
	Phe	vai	50E	THE	rre	Asp	Leu		Asn	Tyr	Gly	Cys	Thr	Ile	Pro	Glu
	lve	ፖ ኬ ո	595 San	0	•		_	600					605			
50	Lys	610	ser	cys	Ser	Val		Gly	Trp	Gly	Tyr	Thr	Gly	Leu	Ile	Asn
			01	•			615					620				
	Tyr .	nsp	ory	reu			Val	Ala	His	Leu	Tyr	Ile	Met	Gly	Asn	Glu
	625	O	٠.	<u> </u>		630					635					640
55	Lys (∨ys :	ser			His	Arg	Gly	Lys		Thr	Leu	Asn	Glu	Ser	G1u
					645					650					655	

		~ ,_		660	nia	ara	Lys	116		ser	GIY	Pro	Cys		GIA	Asp	
	_			660					665					670			
	Tyr	Gly		Pro	Leu	Val	Cys	Glu	Gln	His	Lys	Met	Arg	Met	Val	Leu	
10			675					680					685				
10	Gly	Val	Ile	Val	Pro	Gly	Arg	Gly	Cys	Ala	Ile	Pro	Asn	Arg	Pro	Gly	
		690	•				695					700					
	Ile	Phe	Va1	Arg.	Val	Ala	Tyr	Tyr	Ala	Lys	Trp	Ile	His	Lvs	Ile	Ile	
٠.	705					710	•	-		•	715			•		720	
15			Tvr	Lys	Vel	•	Gln	Son			,-,					,-0	
		••••	-3-	_			OIII	Je.									
					725	•											•
	(2) INFO	RMAT	TON 1	FOR S	EO 1	אר ס	1. 5.										
20	(-) 200			· OIL C			٠.).	•									
	(i)	SEQ	UENC	Е СНА	RACT	reris	TICS	S:									
,				NGTH:					3								
				PE: n			_										
25		(C) ST	RANDE	DNE	SS: d	loub.	le									
2.5		(D) TO	POLOC	Y: :	linea	ar										
	(ii)	MOL	ECUL	E TYF	E: I	DNA	gen	omic)								
	12221	uvo	OTUT	TTO A I													
30	(iii)	піг	Oine.	IICAL	,: Y:	: >> .	. '										
	(iii)	ANT	I-SE	NSE:	NO												
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	(xi)	SEQ	UENC	E DES	CRI	PTIO	۷: SI	EQ I	D NO	: 5:							
35																	
	ATGTGGGT	GA C	CAAA	CTCCI	GÇ	CAGC	CCTG	CTG	CTGC.	AGC	ATGT	CCTC	CT G	CATC	TCCT	C.	60
	CTGCTCCC	CA T	CGCC.	ATCC	CT	ATGC	AGAG	GGA	CAAA	GGA	AAAG	AAGA.	AA T	ACAA	TTCA	T	120
	GAATTCAA	AA A	ATCA	GCAAA	GA	CTAC	CCTA	ATC	AAAA'	TAG	ATCC	AGCA	CT G	AAGA	TAAA	A	180
40	ACCAAAAA	AG T	GAAT	ACTG	AG	ACCA	ATGT	GCT	AATA	GAT	GTAC	TAGG	AA T	AAAG	GACT	Т	240
•	CCATTCAC																300
	TTCAATAG																360
	AACAAAGA																420
45	ACCITATEAC																480
	AGCTTTTT CGAGGGGA																540 600
	TGTGACAT																660
	GGTCTCAT																720
	CACCGGCA																780
50	CGCAATCC																840
	GAGTACTG																900
	GAAACAAC																960
	TGGAATGG																1020
55																	

	CCTGAAAATT TCAAGTGCAA GGACCTACGA GAAAATTACT GCCGAAATCC AGATGGGTCT	
5	GAATCACCCT GGTGTTTTAC CACTGATCCA AACATCCGAG TTGGCTACTG CTCCCAAATT	1080
	CCAAACTGTG ATATGTCACA TGGACAAGAT TGTTATCGTG GGAATGGCAA AAATTATATG	1140
	GGCAACTTAT CCCAAACAAG ATCTGGACTA ACATGTTCAA TGTGGGACAA GAACATGGAA	1200
	GACTTACATC GTCATATCTT CTGGGAACCA GATGCAAGTA AGCTGAATGA GAATTACTGC	1260
10	CGAAATCCAG ATGATGATGC TCATGGACCC TGGTGCTACA CGGGAAATCC ACTCATTCCT	1320
10	TGGGATTATT GCCCTATTTC TCGTTGTGAA GGTGATACCA CACCTACAAT AGTCAATTTA	1380
	GACCATCCCG TAATATCTTG TGCCAAAACG AAACAATTGC GAGTTGTAAA TGGGATTCCA	1440
	ACACGAACAA ACATAGGATG GATGGTTAGT TTGAGATACA GAAATAAACA TATCTGCGGA	1500
	GGATCATTGA TAAAGGAGAG TTGGGTTCTT ACTGCACGAC AGTGTTTCCC TTCTCGAGAC	1560
15	TTGAAAGATT ATGAAGCTTG GCTTGGAATT CATGATGTCC ACGGAAGAGG AGATGAGAAA	1620
	TGCAAACAGG TTCTCAATGT TTCCCAGCTG GTATATGGCC CTGAAGGATC AGATCAGAAA	1680
	TTAATGAAGC TTGCCAGGCC TGCTGTCCTG GATGATTTTG TTAGTACGAT TGATTTACCT	1740 1800
	AATTATGGAT GCACAATTCC TGAAAAGACC AGTTGCAGTG TTTATGGCTG GGGCTACACT	
20	GGATTGATCA ACTATGATGG CCTATTACGA GTGGCACATC TCTATATAAT GGGAAATGAG	1860
	AAATGCAGCC AGCATCATCG AGGGAAGGTG ACTCTGAATG AGTCTGAAAT ATGTGCTGGG	1920
	GCTGAAAAGA TTGGATCAGG ACCATGTGAG GGGGATTATG GTGGCCCACT TGTTTGTGAG	1980
	CAACATAAAA TGAGAATGGT TCTTGGTGTC ATTGTTCCTG GTCGTGGATG TGCCATTCCA	2040 2100
25	AAICGICCIG GTATTTTTGT CCGAGTAGCA TATTATGCAA AATGGATACA CAAAATTATT	2160
25	TTAACATATA AGGTACCACA GTCATAG	2187
30	(2) INFORMATION FOR SEQ ID NO: 6: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2576 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
35	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: mRNA	
40	(iii) HYPOTHETICAL: YES	
	(iii) ANTI-SENSE: NO	
	(ix) FEATURE:	
45	(A) NAME/KEY: CDS	
	(B) LOCATION: join(1022285, 22892294, 22982336, 23402384, 23882480, 24842507, 25142522, 25262570)	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
	GGGCUCAGAG COGACUGGCU CUUUUAGGCA CUGACUCCGA ACAGGAUUCU UUCACCCAGG	60

	CAU	CUCCI	UCC .	AGAG	GGAU	CC G	CCAG	CCCGI	J CC	AGCA	GCAC	C A	UG U	GG GI	JG A	CC	113
5						•						Me		rp Va	al Ti	ır	
													1				
		CUC															161
10		Leu	Leu	Pro	Ala		Leu	Leu	Gln	His		Leu	Leu	His	Leu		
	5					10					15					20	
	CUG	CUC	ccc	AUC	GCC	AUC	ccc	UAU	GCA	GAG	GGA	CAA	AGG	AAA	AGA	AGA	209
	Leu	Leu	Pro	Ile	Ala	Ile	Pro	Tyr	Ala	Glu	Gly	Gln	Arg	Lys	Arg	Arg	
15					25					30					35		
	AAU	ACA	AUU	CAU	GAA	UUC	AAA	AAA	UCA	GCA	AAG	ACU	ACC	ĊUA	AUC	AAA	_257
		Thr															
20				40					45					50			
	AIIA	GAU	CCA	GCA	CUG	AAC	Δ11Δ	A A A	٨٥٥	A A A		CUC	114.4	ACII	CCA	CAC	205
		Asp															305
			55			·		60		•			65				
25																	
		UGU															353
	OIII	Cys 70	UTG	USII	urR	cys	75	ALR	ASII	Lys	GLÅ	B0	Pro	rne	Inr	Cys	
30		•					.,					•					
		GCU															401
	Lys 85	Ala	Phe	Val	Phe		Lys	Ala	Arg	Lys		Cys	Leu	Trp	Phe		
	O)					90					95		•		,	100	
35	UUC	AAU	AGC	AUG	UCA	AGU	GGA	GUG	AAA	AAA	GAA	บบบ	GGC	CAU	GAA	ບບບ	449
	Phe	Asn	Ser	Met	Ser	Ser	Gly	Val	Lys	Lys	Glu	Phe	Gly	His	Glu	Phe	
					105					110					115		
40	GAC	CUC	UAU	GAA	AAC	AAA	GAC	UAC	AUU	AGA	AAC	UGC	AUC	AUU	GGU	AAA	497
		Leu															
				120					125					130			
40	GGA	CGC	ACC	1140	AAC	CCA	۸۵۸	CIIA	иси	AUC	ACU	440	ACU	ccc	MIC		545
45		Arg															כדכ
			135	-	•	•		140					145	•			
															•		
50	_	CAG															593
	oys	Gln 150	rro	ırp	ser	ser	Met 155	тте	Pro	HIS	GLU	160	ser	rne	Leu	Pro	

5	UCG Ser 165	ser	UAU Tyr	CGG Arg	GCU	AAA Lys 170	GAC Asp	CUA Leu	CAG Gln	GAA Glu	AAC Asn 175	Tyr	UGU Cys	CGA Arg	AAU Asn	CCU Pro 180	641
10	CGA Arg	GGG Gly	GAA Glu	GAA Glu	GGG Gly 185	GGA Gly	CCC Pro	UGG Trp	UGU Cys	UUC Phe 190	Thr	AGC Ser	AAU Asn	CCA Pro	GAG Glu 195	GUA Val	689
15	CGC Arg	UAC Tyr	GAA Glu	GUC Val 200	UGU Cys	GAC Asp	AUU Ile	CCU Pro	CAG Gln 205	UGU Cys	UCA Ser	GAA Glu	GUU Val	GAA Glu 210	UGC Cys	AUG Met	737
20	ACC Thr	UGC Cys	AAU Asn 215	GGG Gly	GAG Glu	AGU Ser	UAU Tyr	CGA Arg 220	GGU Gly	CUC Leu	AUG Met	GAU Asp	CAU His 225	ACA Thr	GAA Glu	UCA Ser	<i>]</i> 85
25	GGC Gly	AAG Lys 230	AUU Ile	UGU Cys	CAG Gln	CGC Arg	UGG Trp 235	GAU Asp	CAU His	CAG Gln	ACA Thr	CCA Pro 240	CAC His	CGG Arg	CAC His	AAA Lys	833
30	UUC Phe 245	UUG Leu	CCU Pro	GAA Glu	AGA Arg	UAU Tyr 250	CCC Pro	GAC Asp	AAG Lys	GGC Gly	บบบ Phe 255	GAU Asp	GAU Asp	AAU Asn	UAU Tyr	UGC Cys 260	881
35	CGC Arg	AAU Asn	CCC Pro	GAU Asp	GGC Gly 265	CAG Gln	CCG Pro	AGG Arg	CCA Pro	UGG Trp 270	UGC Cys	UAU Tyr	ACU Thr	CUU Leu	GAC Asp 275	CCU Pro	929
	CAC His	ACC Thr	CGC Arg	UGG Trp 280	GAG Glu	UAC Tyr	UGU Cys	Ala	AUU Ile 285	AAA Lys	ACA Thr	UGC Cys	GCU Ala	GAC Asp 290	AAU Asn	ACU Thr	977
40	AUG Met	AAU Asn	GAC Asp 295	ACU Thr	GAU Asp	GUU Val	CCU Pro	UUG Leu 300	GAA Glu	ACA Thr	ACU Thr	GAA Glu	UGC Cys 305	AUC Ile	CAA Gln	GGU Gly	1025
45	CAA Gln	GGA Gly 310	GAA Glu	GGC Gly	UAC Tyr	Arg	GGC Gly 315	ACU Thr	GUC Val	AAU Asn	ACC Thr	AUU Ile 320	UGG Trp	AAU Asn	GGA Gly	AUU Ile	1073
50	CCA Pro 325				Trp												1121

5	CCU	GAA	AAU	UUC	AAG	UGC	AAG	GAC	CUA	CGA	GAA	AAU	HAC	HGC	CGA	AAU	1160
	Pro	Glu	Asn	Phe	Lys 345	Cys	Lys	Asp	Leu	Arg 350	Glu	Asn	Tyr	Cys	Arg 355	Asn	1169
	CCA	GAU	GGG	UCU	GAA	UCA	CCC	UGG	UGH	шш	ACC	ACU	CALL	CCA	140	AUC	1217
10	Pro	Asp	Gly	Ser 360	Glu	Ser	Pro	Trp	Cys 365	Phe	Thr	Thr	Asp	Pro 370	Asn	Ile	1217
	CGA	GUU	GGC	UAC	UGC	UCC	CAA	AUU	CCA	AAC	UGU	GAU	Alig	IICA	CAII	CCA	1265
15	Arg	Val	Gly 375	Туг	Cys	Ser	Gln	Ile 380	Pro	Asn	Cys	Asp	Met 385	Ser	His	Gly	1209
	CAA	GAU	UGU	UAU	CGU	GGG	AAU	GGC	AAA	AAU	UAU	AUG	GGC	AAC	UUA	UCC	1313
20	Cln	Asp 390	Cys	Tyr	Arg	Gly	Asn 395	Gly	Lys	Asn	Tyr	Met 400	Gly	Asn	Leu	Ser	-5-3
	CAA	ACA	AGA	UCU	GGA	CUA	ACA	UGU	UCA	AUG	UGG	GAC	AAG	AAC	AUG	GAA	1361
	Gln	Thr	Arg	Ser	Gly	Leu	Thr	Cys	Ser	Met	Trp	Asp	Lys	Asn	Met	Glu	1,01
25	405					410					415			•		420	
	GAC	UUA	CAU	CGU	CAU	AUC	UUC	UGG	GAA	CCA	GAU	GCA	AGU	AAG	CUG	AAU	1409
	Asp	Leu	His	Arg	His 425	Ile	Phe	Trp	Glu	Pro 430	Asp	Ala	Ser	Lys	Leu 435	Asn	2,07
30	GAG	ΔΔΙΙ	1140	ucc	CCA	A A T 1	CCA	CATI	C4**								
			Tyr									CAU His					1457
35	UAC	ACG	GGA	AAU	CCA	CUC	IIUA	CCII	UGG	GAU	HAII	UGC	CCII	ΔΙΙΙΙ	ucu	CCU	1505
		Thr										Cys					1505
40	UGU	GAA	GGU	GAU	ACĊ	ACA	CCU	ACA	AUA	GUC	AAU	UUA	GAC	CAU	CCC	GUA	1553
	Cys					Thr						Leu 480					-)))
	AUA.	UCU	UGU	GCC	AAA	ACG	AAA	CAA	UUG	CGA	GUU	GUA	AAU	GGG	AUII	CCA	1601
45												Val					1001
	485					490					495			-		500	
	ACA	CGA	ACA	AAC	AUA	GGA	UGG	AliG	GUU	AGU	шс	ÄGA	ዘልሮ	ΔCA	ΔΔΙΙ	ΔΔΔ	1600
50												Arg					1649
					505	•	•			510			- J -		515	-, 0	

5	CAU	AUC	UGC	GGA	GGA	UCA	UUG	AUA	AAG	GAG	AGU	UGG	GUU	CUU	ACII	GCA	1697
	His	Ile	Cys	Gly 520	Gly	Ser	Leu	Ile	Lys 525	Glu	Ser	Trp	Val	Leu 530	Thr	Ala	10)1
10	CGA Arg	CAG Gln	UGU Cys 535	UUC Phe	CCU Pro	UCU Ser	CGA Arg	GAC Asp 540	UUG Leu	AAA Lys	GAU Asp	UAU Tyr	GAA Glu 545	GCU Ala	UGG Trp	CUU Leu	1745
15	GGA Gly	AUU Ile 550	CAU His	GAU Asp	GUC Val	CAC His	GGA Gly 555	AGA Arg	GGA Gly	GAU Asp	GAG Glu	AAA Lys 560	UGC Cys	AAA Lys	CAG Gln	GUU Val	1793
20						CUG Leu 570											1841
25						AGG Arg											1889
30						UAU Tyr											1937
						GGC Gly											1985
35						CUC Leu											2033
40						GUG Val 650											2081
4 5						UCA Ser											2129
50						CAU His											2177

5	CCU	GGU	CGU	GGA	UGU	GCC	AUU	CCA	AAU	CGU	CCU	GGU	AUU	w	GUC	CGA	2225
	Pro	Gly		Gly	Cys	Ala	Ile	Pro	Asn	Arg	Pro	Gly	Ile	Phe	Val	Arg	
			695					700					705				
	GIJA	GCA	11411	11411	CCA	A A A	ucc	AUA	CAC	A A A	ATTIT	A 1 171 1	7777.4		****		2222
10								Ile									2273
		710	-3-	.,.		LJS	715	116	1112	Lys	116	720	Leu	Int	ıyı.	Lys	
		•					,					,					
	GUA	CCA	CAG	UCA	UAG	CUG	AAG	UAA	GUG	UGU	CUG	AAG	CAC	CCA	CCA	AUA	2321
15	Val	Pro	Gln	Ser		Leu	Lys		Val	Cys	Leu	Lys	His	Pro	Pro	Ile	
	725		•				730						735				
	САА	CUG	IICII	1001	ΔCΔ	HGA	AGA	บบบ	CAC	A.C.A	ATIC	IICC	A A 11	TITTA	4 4 4	11011	2260
		Leu				UGA		Phe									2369
20		740						745		0		P	750	LCu	D , 5	0,5	
	CAC	UUA	CAA	CAA	UCC	UAA	GAC	AAC	UAC	UGG	AGA	GUC	AUG	UUU	GUU	GAA	2417
	His	Leu	G1n	Gln	Ser		Asp	Asn	Tyr	Trp	Arg	Val	_	Phe	Val	Glu	
25		755						760					765				
	IIIIA	CHC	ATILL	1144	GIIII	IIAII	GGG	UGU	11171	CUG	IIIIG	וחחו	nen	IIIIG	IICA	CIIC	2465
								Cys									2405
		770				•	775					780	-,-				
30																	
						UGA		GAA									2507
		Phe	Cys	Gln	Cys			Glu	Leu	Arg	Tyr		Gln	Val			
	785						790			•		795					
35	UAAI	JAA (CAU	AUC I	icc i	IGA A	AGA I	UAC 1	IIIG A	AAU (GGA 1	HIA	AAA .	AAA 1	CAC	ACA	2555
			lis :					Tyr I									-222
				{	800						805			-		810	
40					GGA	UGAU	JAA										2576
	Gly	Ile	Phe	Ala													
					815												
45	(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:	7:								
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50					YPE:												
			()	וו נט	OPOL	JUI:	1110	ear									
		(ii) MO	LEGU	LE T	YPE:	pro	tein									

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

5	4	, iti	י אאי	Linr			Leu	Pro	Ala	Leu	Leu	Leu	Gln	His	Val	Leu
	. 1				5					10					15	
	Leu	His	. Leu	Leu	Leu	Leu	Pro	Ile	Ala	Ile	Pro	Tyr	Ala	Glu	Gly	Gln
40				20					25					30		
10	Arg	Lys	Arg	Arg	Asn	Thr	Ile	His	Glu	Phe	Lys	Lys	Ser	Ala	Lys	Thr
			35					40					45			
	Thr	Leu	Ile	Lys	Ile	Asp	Pro	Ala	Leu	Lys	Ile	Lys	Thr	Lys	Lys	Val
15		50)				55					60				
	Asn	Thr	Ala	Asp	Gln	Cys	Ala	Asn	Arg	Cys	Thr	Arg	Asn	Lys	Gly	Leu
	65					70					75					80
20	Pro	Phe	Thr	Cys	Lys	Ala	Phe	Val	Phe	Asp	Lys	Ala	Arg	Lys	Gln	Cys
20					85					90					95	
	Leu	Trp	Phe	Pro	Phe	Asn	Ser	Met	Ser	Ser	Gly	Val	Lys	Lys	Glu	Phe
				100					105					110		
25	Gly	His	Glu	Phe	Asp	Leu	Tyr	Glu	Asn	Lys	Asp	Tyr	Ile	Arg	Asn	Cys
			115					120					125			
	Ile	Ile	Gly	Lys	Gly	Arg	Ser	Tyr	Lys	Gly	Thr	Val	Ser	Ile	Thr	Lys
30		130					135					140				
	Ser	Gly	Ile	Lys	Cys	Gln	Pro	Trp	Ser	Ser	Met	Ile	Pro	His	Glu	His
	145					150					155					160
	Ser	Phe	Leu	Pro	Ser	Ser	Туг	Arg	Gly	Lys	Asp	Leu	Gln	Glu	Asn	Tyr
35					165					170					175	
	Cys	Arg	Asn	Pro	Arg	Gly	Glu	Glu	Gly	Gly	Pro	Trp	Cys	Phe	Thr	Ser
				180					185					190		
40	Asn	Pro	Glu	Val	Arg	Tyr	Glu	Val	Cys	Asp	Ile	Pro	Gln	Cys	Ser	Glu
			195					200					205			
	Val	Glu	Cys	Met	Thr	Cys	Asn	Gly	Glu	Ser	Tyr	Arg	Gly	Leu	Met	Asp
		210					215					220				
4 5	His	Thr	Glu	Ser	Gly	Lys	Ile	Cys	Gln	Arg	Trp	Asp	His	Gln	Thr	Pro
	225					230					235					240
	His	Arg	His	Lys	Phe	Leu	Pro	Glu	Arg	Tyr	Pro	Asp	Lys	Gly	Phe	
50					245					250		-	-		255	•
	Asp	Asn	Tyr	Cys	Arg	Asn	Pro	Asp	Gly		Pro	Arg	Pro	Tro		Tvr
			_	260					265			J	-	270		-5-
	Thr	Leu	Asp	Pro	His	Thr	Arg			Tyr	Cys	Ala	Ile		Thr	Cve
55			275					280	-	- 🗸 -	- J —		285	_, _		-,,.,

	Ala	Asp	Asn	Thr	Met	Asn	Asp	Thr	Asp	Val	Pro	Leu	Glu	Thr	Thr	Glu
		290					295					300				
5	Cys	Ile	Gln	Gly	Gln	Gly	Glu	Gly	Tyr	Arg	Gly	Thr	Val	Asn	Thr	Ile
	305					310					315					320
	Trp	Asn	Gly	Ile	Pro	Cys	Gln	Arg	Trp	Asp	Ser	Gln	Tyr	Pro	His	Glu
10					325					330					335	
	His	Asp	Met	Thr	Pro	Glu	Asn	Phe	Lys	Cys	Lys	qsA	Leu	Arg	Glu	Asn
	_	_		340				•	345					350		
	Tyr	Cys		Asn	Pro	Asp	Gly	Ser	Glu	Ser	Pro	Trp	Cys	Phe	Thr	Thr
15		_	355					360					365			
	Asp		Asn	Ile	Arg	Val	Gly	Tyr	Cys	Ser	Gln	Ile	Pro	Asn	Cys	Asp
		370					375				•	380				•
20	Met	Ser	His	Gly	Gln	Asp	Cys	Tyr	Arg	Gly	Asn	Gly	Lys	Asn	Tyr	Met
	385			_		390					395					400
	GIY	Asn	Leu	Ser	Gln	Thr	Arg	Ser	Gly		Thr	Cys	Ser	Met	Trp	Asp
25	ī	۸	16- A	03	405	_				410				•	415	
	Lys	ASN	met		Asp	Leu	His	Arg		Ile	Phe	Trp	Glu	Pro	Asp	Ala
	San		7	420	0 1		_	_	425					430		
	Ser	Lys		Asn	Glu	Asn	Tyr		Arg	Asn	Pro	Asp		Asp	Ala	His
30	Glv	Pno	435	Cua	T	77%	01	440				_	445			
	ury	450	тър	Cys	Tyr	Inr		Asn	Pro	Leu	lle		Trp	Asp	Tyr	Cys
-	Pro	-	Sar	Aria	Cvc	C1	455	۸		m.	_	460				
35	465	110	Ser	urg	Cys	470	gry	ASP	inr	Inr		Thr	lle	Val	Asn	
		His	Pro	Val	Tlo	•	Cuc	۸1.	1	Тъ.,	475	01				480
		0			Ile 485	561	∪y5	ura	Lys	490	Lys	GIN	Leu	Arg		Vai
40	Asn	G1v	Ile	Pro	Thr	Aro	Thr	Aen	T10		Tnn	No.	17-1	C	495	
				500		8	****	กรแ	505	GI.y	Trp	met	vai		Leu	Arg
	Tyr	Arg	Asn	-	His	Tle	Cvs	G1 v	_	San	1 011	T10	1	510	C	Т
	•	-0	515	-5 -			V	520	dry	Ser	rea	TIE		GIU	ser	1rp
45	Val	Leu		Ala	Arg	Gln	Cvs		Pro	San	Arc	Acn	525	1	۸	Т
		530			****	44. .	535	- 110	110	Ser	urg	540	rea	гàг	кѕр	lyr
			Tro	Leu	Gly	Tle		Asn	Va1	Hic	Glv		C1	Aan	C1	I
50	545				3	550		цор	Val	1113	555	urg	GIŞ	nsp	GIU	_
		Lvs	Gln	Val	Leu		Val	Ser	Gln	Len		Tun	G1	Dno	C1	560
	-	-	•		565				-411	570	• 41	·yı	-1y	110		ary
:	Ser	Asp	Leu	Val	Leu	Met	Lvs	Len	Ala		Pro	ΑΙα	Va1	l eu	575	Δ c =
55		-		580			. -		585	0			·ul	590	nap	vah
				-					/					,,,,		

	Phe	Val	Ser	Thr	Ile	Asp	Leu	Pro	Asn	Tyr	Gly	Cys	Thr	· Ile	Pro	Glu
			595					600					605			
5	Lys	Thr	Ser	Cys	Ser	Val	Tyr	Gly	Trp	Gly	Tyr	Thr	Gly	Leu	Ile	Asn
		610					615					620				
	Tyr	Asp	Gly	Leu	Leu	Arg	Val	Ala	His	Leu	Tyr	Ile	Met	Gly	Asn	Glu
10	625	i				630					635					640
	Lys	Cys	Ser	Gln	His	His	Arg	Gly	Lys	Val	Thr	Leu	Asn	Glu	Ser	Glu
					645					650					655	
4.00	Ile	Cys	Ala	Gly	Ala	Glu	Lys	Ile	Gly	Ser	Gly	Pro	Cys	Glu	Gly	Asp
15				660					665					670		
	Tyr	Gly	Gly	Pro	Leu	Val	Cys	Glu	Gln	His	Lys	Met	Arg	Met	Val	Leu
			675					680					685			
20	Gly	Val	Ile	Val	Pro	Gly	Arg	Gly	Cys	Ala	Ile	Pro	Asn	Arg	Pro	Gly
		6 <u>9</u> 0					695					700				
	Ile	Phe	Val	Arg	Val	Ala	Tyr	Tyr	Ala	Lys	Trp	Ile	His	Lys	Ile	Ile
25	705					710					715					720
	Leu	Thr	Tyr	Lys	Val	Pro	Gln	Ser	Leu	Lys	Val	Cys	Leu	Lys	His	Pro
					725					730					735	
	Pro	Ile	Gln		Ser	Phe	Thr	Arg	Phe	Gln	Arg	Met	Trp	Asn	Leu	Lys
30				740					745					750		
	Cys	His		Gln	Gln	Ser	Asp	Asn	Tyr	Trp	Arg	Val	Met	Phe	Val	Glu
			755					760					765			
35	Ile	Leu	Ile	Asn	Val	Tyr	Gly	Cys	Phe	Leu	Leu	Phe	Cys	Leu	Ser	Val
	_	770					775					780				
		Phe	Cys	Gln	Cys	Ser	Glu	Leu	Arg	Tyr	Met	Gln	Val	His	Ile	Ser
10	785					790					795					800
ю	Arg	Tyr	Leu			Leu	Lys	Lys	His	Thr	Gly	Ile	Phe	Ala	Gly	
					805					810					815	

Claims

20

5 1. A single chain protein selectively enhancing the growth of vascular endothelial cells, characterized in that it comprises the following peptide chains:

(SEQ. ID No. : 1) Arg Asn Thr Ile His Glu Phe 10 1 5 (SEQ. ID No. : 2) Glu Phe Gly His Glu Phe Asp Leu Tyr Glu 15 5 10 1 (SEQ. ID No. : 3) Glu Ser Xaa Val Leu Thr Ala Arg Gln Xaa Phe Pro Ser Arg Asp Leu 5 10 15

and in that it has a molecular weight of from 72,000 to 80,000 Da when determined by SDS polyacrylamide gel electrophoresis or from 79,000 to 85,000 Da when determined under reducing conditions.

- 2. A process for producing the protein according to claim 1 which comprises purifying a serum-free culture supernatant of said human ovarian tumor established cell line, HUOCA-II or HUOCA-III, by combining purification techniques including (a) cation exchange chromatography, (b) heparin affinity chromatography, (c) heparin affinity high performance liquid chromatography and (d) reverse phase high performance liquid chromatography.
- A protein of human origin which contains an amino acid sequence or a portion of the amino acid sequence represented by the following sequence (SEQ ID No.: 4):

Met Trp Val Thr Lys Leu Leu Pro Ala Leu Leu Leu Gln His Val

1 10

Leu Leu His Leu Leu Leu Leu Pro Ile Ala Ile Pro Tyr Ala Glu
20

Gly Gln Arg Lys Arg Arg Asn Thr Ile His Glu Phe Lys Lys Ser
40

Ala Lys Thr Thr Leu Ile Lys Ile Asp Pro Ala Leu Lys Ile Lys
50

50

45

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Thr Lys Lys Val Asn Thr Ala Asp Gin Cys Ala Asn Arg Cys Thr
      Arg Asn Lys Gly Leu Pro Phe Thr Cys Lys Ala Phe Val Phe Asp
      Lys Ala Arg Lys Gln Cys Leu Trp Phe Pro Phe Asn Ser Met Ser
      Ser Gly Val Lys Lys Glu Phe Gly His Glu Phe Asp Leu Tyr Glu
                      110
      Asn Lys Asp Tyr Ile Arg Asn Cys Ile Ile Gly Lys Gly Arg Ser
 10
     Tyr Lys Gly Thr Val Ser Ile Thr Lys Ser Gly Ile Lys Cys Gln
         Trp Ser Ser Met Ile Pro His Glu His Ser Phe Leu Pro Ser
     Ser Tyr Arg Gly Lys Asp Leu Gln Glu Asn Tyr Cys Arg Asn Pro
 15
                      170
     Arg Gly Glu Glu Gly Gly Pro Trp Cys Phe Thr Ser Asn Pro Glu
     Val Arg Tyr Glu Val Cys Asp Ile Pro Gln Cys Ser Glu Val Glu
20
     Cys Met Thr Cys Asn Gly Glu Ser Tyr Arg Gly Leu Met Asp His
     Thr Glu Ser Gly Lys Ile Cys Gln Arg Trp Asp His Gln Thr Pro
                      230
     His Arg His Lys Phe Leu Pro Glu Arg Tyr Pro Asp Lys Gly Phe
25
     Asp Asp Asn Tyr Cys Arg Asn Pro Asp Gly Gln Pro Arg Pro Trp
                      260
     Cys Tyr Thr Leu Asp Pro His Thr Arg Trp Glu Tyr Cys Ala Ile
                                          280
     Lys Thr Cys Ala Asp Asn Thr Met Asn Asp Thr Asp Val Pro Leu
30
                     290
     Glu Thr Thr Glu Cys lie Gln Gly Gln Gly Glu Gly Tyr Arg Gly
     Thr Val Asm Thr Ile Trp Asm Gly Ile Pro Cys Glm Arg Trp Asp
35
                     320
     Ser Gln Tyr Pro His Glu His Asp Met Thr Pro Glu Asn Phe Lys
     Cys Lys Asp Leu Arg Glu Asn Tyr Cys Arg Asn Pro Asp Gly Ser
                     350
40
     Glu Ser Pro Trp Cys Phe Thr Thr Asp Pro Asn Ile Arg Val Gly
     Tyr Cys Ser Gln Ile Pro Asn Cys Asp Met Ser His Gly Gln Asp
                     380
    Cys Tyr Arg Gly Asn Gly Lys Asn Tyr Met Gly Asn Leu Ser Gln
45
    Thr Arg Ser Gly Leu Thr Cys Ser Met Trp Asp Lys Asn Met Glu
                     410
    Asp Leu His Arg His Ile Phe Trp Glu Pro Asp Ala Ser Lys Leu
    Asn Glu Asn Tyr Cys Arg Asn Pro Asp Asp Asp Ala His Gly Pro
50
                     440
    Trp Cys Tyr Thr Gly Asn Pro Leu Ile Pro Trp Asp Tyr Cys Pro
```

```
The Ser Arg Cys Glu Gly Asp Thr Thr Pro Thr He Val Asn Leu
    Asp His Pro Val Ile Ser Cys Ala Lys Thr Lys Gln Leu Arg Val
                                          490
    Val Ash Cly Ile Pro Thr Arg Thr Ash Ile Gly Trp Met Val Ser
    Leu Arg Tyr Arg Asn Lys His Ile Cys Gly Gly Ser Leu Ile Lys
    Glu Ser Trp Val Leu Thr Ala Arg Gln Cys Phe Pro Ser Arg Asp
10
                     530
    Leu Lys Asp Tyr Glu Ala Trp Leu Gly Ile His Asp Val His Gly
                                         550
    Arg Gly Asp Glu Lys Cys Lys Gln Val Leu Asn Val Ser Gln Leu
                     560
    Val Tyr Gly Pro Glu Gly Ser Asp Leu Val Leu Met Lys Leu Ala
15
                                         580
    Arg Pro Ala Val Leu Asp Asp Phe Val Ser Thr Ile Asp Leu Pro
                     590
    Asn Tyr Gly Cys Thr Ile Pro Glu Lys Thr Ser Cys Ser Val Tyr
    Gly Trp Gly Tyr Thr Gly Leu Ile Asn Tyr Asp Gly Leu Leu Arg
                     520
    Val Ala His Leu Tyr Ile Met Gly Asn Glu Lys Cys Ser Gln His
                                         64C
    His Arg Gly Lys Val Thr Leu Ash Glu Ser Glu Ile Cys Ala Cly
                     650
25
    Ala Glu Lys Ile Gly Ser Gly Pro Cys Glu Gly Asp Tyr Gly Gly
    Pro Leu Val Cys Glu Gln His Lys Met Arg Met Val Leu Gly Val
                                                              690
                     680
    Ile Val Pro Gly Arg Gly Cys Ala Ile Pro Asn Arg Pro Gly Ile
30
                                         700
    Phe Val Arg Val Ala Tyr Tyr Ala Lys Trp Ile His Lys Ile Ile
                     710
    Leu Thr Tyr Lys Val Pro Gla Ser
```

- 4. A pharmaceutical composition which contains the protein of claim 1 or 3 as an active ingredient.
- A DNA fragment which contains a nucleotide sequence or a portion of the nucleotide sequence below
 (SEQ ID No.: 5):

ATG TGG GTG ACC ARA CTC CTG CCA GCC CTG CTG CTG CAG CAT

GTC CTG CTG CAT CTC CTG CTG CTC CCC ATC GCC ATC CCC TAT

45
GCA GAG GGA CAA AGG AAA AGA AGA AAT ACA ATT CAT GAA TTC

93
AAA AAA TCA GCA AAG ACT ACC CTA ATC AAA ATA GAT CCA GCA

141
CTG AAG ATA AAA ACC AAA AAA GTG AAT ACT GCA GAC CAA TGT

189
GCT AAT AGA TGT ACT AGG AAT AAA GGA CTT CCA TTC ACT TGC

55

		GCT T								205			
	TTC	CCC T	TC AA	r AGC	ATG	TCA	AGT	GGA	GTG	AAA	AAA	GAA	TTT
5	GGC	CAT G	AA TT	r GAC	CTC	TAT	GAA	AAC	AAA	GAC	TAC	333 ATT	AGA
	λAC 381	TGC A	TC AT	GGT	' AAA	GGA	CGC	AGC	TAC	AAG	GGA	AÇA	GTA
10		ATC A	CT AAC	AGT	GGC	ATC	AAA	TGT	CAG	ccc	TGG	AGT	TCC
	ATG	ATA C		GAA	CAC	AGC	TTT	TTG	CCT	TCG	AGC	TAT	CGG
	GGT	AAA G	AC CTA		GAA	AAC	TAC	TGT	CGA	AAT	CCT	CGA	GGG
15	GAA	GAA GO	G GGA	ccc	TGG	TGT	TTC		AGC .	P.A.T	CCA	GAG	GTA
	CGC	TAC GA	A GTC	TGT	GAC	ATT	CCT	573 CλG			GAA	GTT	SAA
	TGC	ATG AC	C TGC	AAT	GGG	GAG	AGT	TAT	CGA (621 3GT			GAT
20	CAT 1	ACA GA	LA TCA	GGC	AAG	ATT	TGT	CAG	CGC 1	rgg	GAT (699 CAT	CAG
	ACA C	CA CA	C CGG	CAC	AAA	TIC	TTG (CCT (BAA <i>H</i>	.GA	TAT (cc (GAC
25		GC TT	r gat										
		76. GG CC	2	TGC									
	GAG T	AC TG	r GCA	813 ATT	AAA I	ACA '	TGC (SCT G	AC A	AT Z	ACT A	.TG #	AAT
30	GAC A	CT GAT	GTT	CCT '	TTG (861 GAA 1	ACA A	ACT G	AA T	GC I	ATC C	AA C	GT
	CAA G	GA GAZ	GGC	TAC 2	AGG (GC 2	ACT G	909 STC A			T TT	GG A	LAT
35	GGA A	TT CCA	TGT	CAG (CGT T	rgg (GAT T	CT C	AG T	57 Ar c		_	:AG
	CAT G	AC ATO	ACT	CCT (SAA A	LAT 3	TTC A	AG T	GC A	AG G	AC C	05 TA C	GA
	GAA A1 1053	AT TAC	TGC	CGA A	AAT C	CA	AT G	GG T	CT G	A T	CA C	CC T	GG
40	TGT T1	77 ACC 1101	ACT (GAT (CCA A	AC A	TC C	GA G	ľፕ GO	SC T	AC TO	GC I	СС
	CAA AT	T CCA	AAC 1	CGT G	AT A	TG I	CA C	AT G	GA CA	AA G	AT T	GT T	AT
	CGT GG	G AAT			AT T	AT A	TG G	GC A	AC TI	'A T	cc ca	AA A	CA
45	AGA TO				'GT T	CA A	12	2 5					
	GAC TT						GG G	AA CC	120	2			
50	CTG AA								I GA	Ψ G	1 7 4	-	
	GGA CC										T TG	G GA	
	TAT TG 1389												
5 5	ATA GT												
	AAA CA	A TTG	CGA G	፻፻ G: 85	TA AA	AT GO	GG AT	T CC	A AC	A CO	A AC	A AF	s.C

	AT.	A GG	A TG	G AT	G GTT	r AGT	TTC		A TAC	AG?	A AA	r aa	A CAS	DTA T
	TG	C GG	A GG	A TC	A TTO	ATA	A AAC	GAC	AGI 1581		G GT	r cr	T AC	r GCA
5	CG	A CA	G TG	T TT	C CC1	r TCI	r CG?	A CAC	TTG	AA	A GA1		r GA	A GCT
	TG	G CT	T GG	A AT	T CAT	GA?	GTC	CAC	GG.	AGA	A GG	A GA	r GAC 167	G AAA 7
10														CCT
	17	25		•										r GCT
	GT	C CT	G GA		r TTI	r GTI	P AGT	ACC	ATI	' GA1	r TT	A CC	r aa:	TAT
15														
	GGA	TGC	ACA		CCT 821	GAA	AAG	ACC	AGT	TGC	AGT	GTT	TAT	GGC
	TGG	GGC	TAC	ACT	GGA		ATC 869	AAC	TAT	GAT	GGC	CTA	TTA	CGA
20			CVI		TAT			1	917			٠		
					AAG					1	.965			
25					AAG							2	2013	
					CTT									
	206:	L			ATT									
30	AAT		CCT 2109	GGT	ATT	TTT	GTC	CGA	GTA	GCA	TAT	TAT	GCA	AAA
	TGG	ATA	CAC		ATT 2157	ATT	TTA	ACA	TAT	AAG	GTA	CCA	CAG	TÇA
	TAG	;	2187											

wherein at least one base may be substituted based on the degeneracy of genetic code.

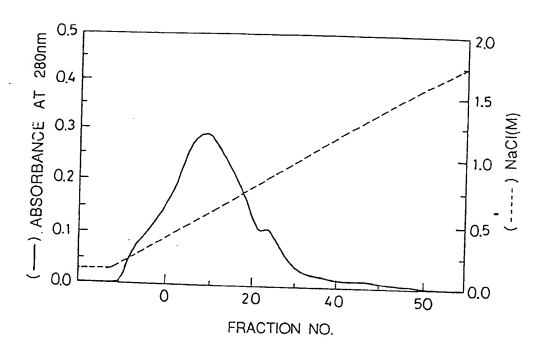
- 6. A single chain protein having an activity to enhance the growth of vascular endothelial cells obtainable from the DNA fragment of claim 5.
- 7. A DNA fragment complementary to the DNA fragment of claim 5.

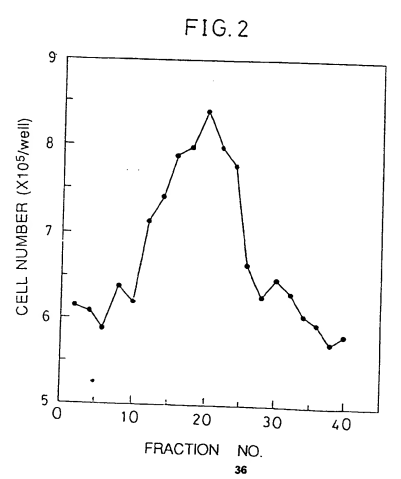
35

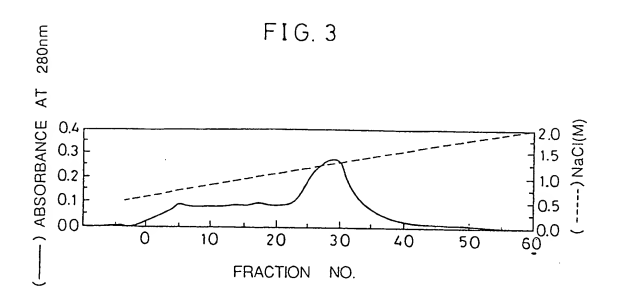
50

- 8. An expression vector which contains the DNA fragment of claim 5.
- 9. A transformant transformed with the DNA fragment of claim 5.
- 10. A transformant transformed with the expression vector of claim 8.

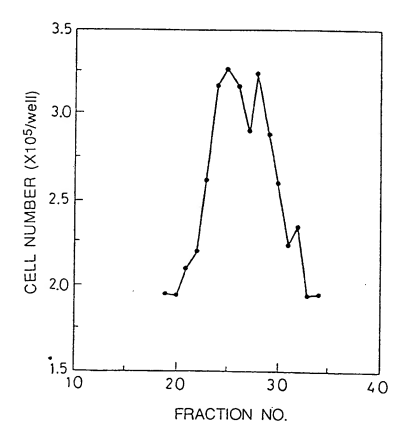
FIG. 1

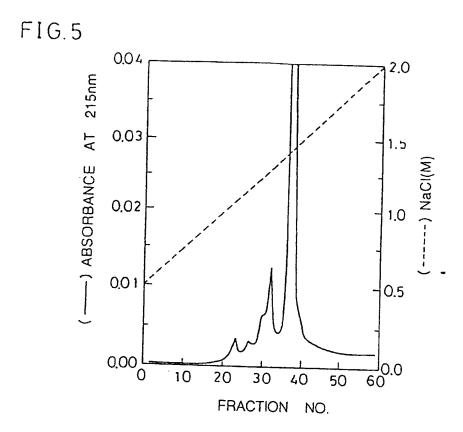






F I G. 4





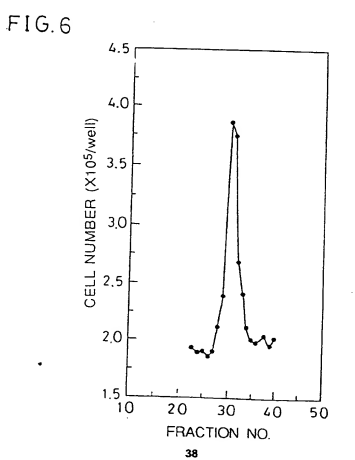


FIG. 7

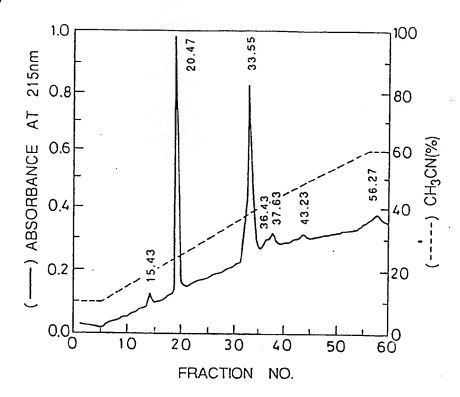
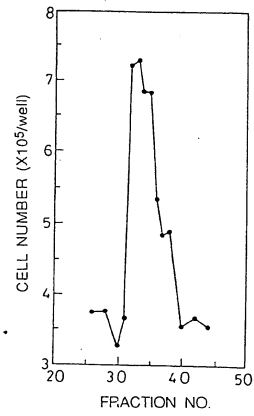
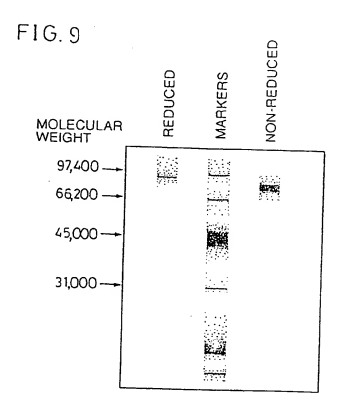
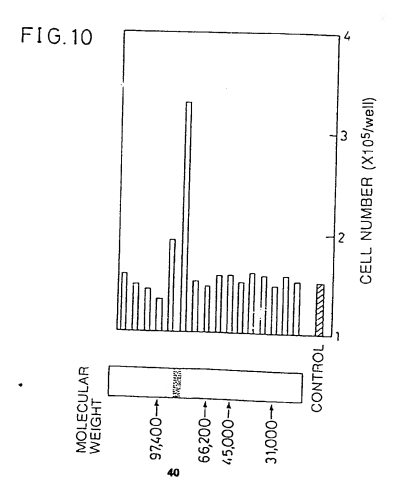


FIG. 8







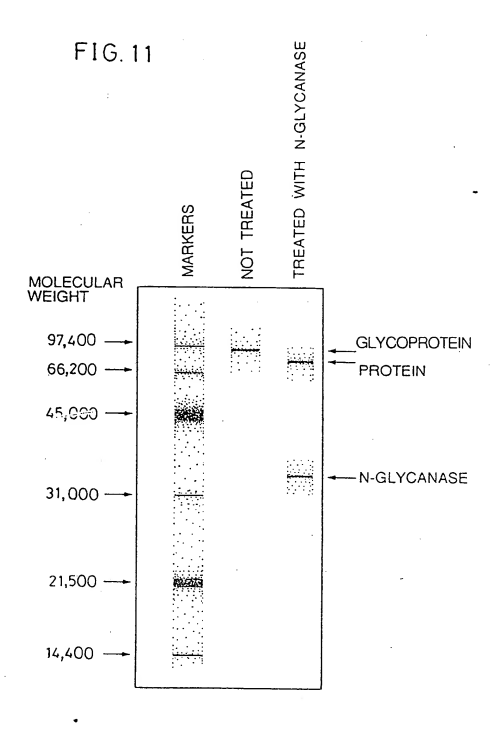


FIG. 12

1	G	G GC	U CA	G AG	ic co	A CU	G GC	ບ ເບ	ָטט ט.	A GG	C AC	U GA	C HC	·C G	4 C	۸.	GAU	
48	UC	บ บบ	C AC	C CA	G GC	A UC	J CC	U CC	A GA	G GG.	A UC	C GC	C AG	c c		AG	GAU	4
1 96			Me	t Tr	n Va	1 Th		- •	_									9
15	Va:	l Le	u Le	n Hi					_									14
144						u Lei C CUC				, 400	- 60	CAU	CC	C UA	.U G	CA	GAG	3 19
31 192						g Arg A AGA				CAL	GA	A UU(AA	A AA	A UC	`A	GCA	4 23
47	Lys	Th	r Thi	r Lei	. T1.	. T	- · -		_									23
240										. GCA	CUC	JAAL	i AUA	. AA	A AC	C	AAA	6 28
63 288	Lys	Va.	l Ası	Th	Ala	A SAC	Gln	Cys	Ala	Asn	Are	Cvs	Thi	- Ar-	or se		7	_
200	AAA	. 600	, AA(J AC	J GC	A GAC	CAA	UCU	GCU	AAU	AGA	UGU	ACI	JAG	G AA	Ü	AAA	7 33
79	Gly	Lei	Pro	Phe	Thr		1											33
336	GGA	CUU	I CCA	UUC	ACU	UGC	AAG	GCU	. Pne ! Don	CUU	Phe	Asp	Lys	Al	a Ar	g	Lys	9
95	Cl.	0		_						300	000	GAU	AAA	GC,	A AG	A	AAA	38
384	CAA	Uys	Leu	Trp	Phe	Pro	Phe	Asn	Ser	- Mail	Ser	Ser	Jly	Va:	1 1.v	•	I.ve	
					_			MAG	AGC	AUG	UCA	AGU	GGA	GUC	AA E	A	AAA	11 43
111	Glu	Phe	Glv	Hic	Clar	Dha		-	_									7.5
432	GAA	UUU	GGC	CAU	GAA	UUU	GAC	CUC	UAU	GAA	ASD	Lys	Asp	Tyr	Il	е,	Arg	12
127	Asn	Cve	Tla	7 1-	~ .					••••	11/10	AAA	GAC	UAC	: AU	U ,	AGA	47
480	AAC	UGC	AUC	TIE	GTA	Lys AAA	Gly	Arg	Ser	Tyr	Lys	Gly	Thr	Val	. Se	r	[]e	14
										UAC	AAG	GGA	ACA	GUA	UC	13 4	2014	52
143 528	Thr	Lys	Ser	Glv	Tlo	T ***	a		_									-
328	ACU	AAG	AGU	GGC	AUC	AAA	UGU	CAG	CCC	UGG	AGU	UCC	AUG	TIE	Pr) }	lis	15
159	Glu	His	Ser	Phe	1 011	D	^	_										57
576	GAA	CAC	AGC	UUU	UUG	Pro CCU	Ser	Ser	Tyr	Arg	Gly	Lys	Asp	Leu	Gli	1 (ilu	17
175									50	CGG	990	AAA	GAC	CUA	CAC	3 (AA	62
624	ASD	Tyr	Cys	Arg	Asn	Pro CCU	Arg	Gly	Glu	Glu	Glv	Glv	Pro	Trn	0			
• • •	7.7.0	UAC	UGU	CGA	AAU	CCU	CGA	GGG	GAA	GAA	GGG	GGA	CCC	UGG	USI	: <u>}</u>	'he	19
191	Thr	Ser	Asn	Pro	Gla	Val		_										67
672	ACA	AGC	AAU	CCA	GAG	Val GUA	CGC	UAC	GIU	Val	Cys	Asp	Ile	Pro	Glr	C	уs	20
207								••••	~·	GOC.	060	GAC	AUU	CCU	CAC	: 17	CII	71
	UCA	GAA	CHIL	Glu	Cys	Met AUG .	Thr	Cys	Asn	Gly	Glu	Ser	Tyr	Arø	Glu			
							_			000	UAG	~60	CAU	CGA	GGH	_	110	22 76
223	net .	ASD	His	Thr	C 1	50-	~	_										, 0
768	AUG	GAU	CÅN	ACA	GAA	UCA (GGC /	AAG .	AUU I	UGU (CAG	Arg	Trp UGG	Asp GAU	His CAU	G C	ln AG	23 81

FIG. 12 (cont.)

239	Thr	Pro	His	Arg	His	Lys	Phe	Leu	Pro	Glu	Arg	Tyr	Pro	Asp	Lys	Gly	25
816		CCA	CAC	CGG	CAC	Aaa	UUC	UUG	CCU	GAA	AGA	UAU	CCC	GAC	aag	GGC	86
255	Phe	Asp	Asp	Asn	Tyr	Cys	Arg	Asn	Pro	Asp	Gly	Gln	Pro	Arg	Pro	Trp	27
864	UUU	GAU	CAU	AAU	UAU	UGC	CGC	AAU	CCC	GAU	GGC	CAG	CCG	AGG	CCA	.UGG	91
271	Cys	Tyr	Thr	Leu	Asp	Pro	His	Thr	Arg	Trp	Glu	Tyr	Cys	Ala	Ile	Lys	28
912	UGC	UAU	ACU	CŲU	GAC	CCU	CAC		CGC	UGG	GAG	UAC	UGU	GCA	AUU	AAA	95
287 960	Thr ACA	Cys	Ala GCU	Asp GAC	Asn AAU	Thr	Met AUG	Asn AAU	Asp GAC	Thr ACU	Asp GAU	Val GUU	Pro CCU	Leu UUG	Glu GAA	Thr ACA	30 100
303 1008	Thr	Glu GAA	Cys UGC	Ile AUC	Gln CAA	Gly GGU	Gln CAA	Gly GGA	Glu GAA	Gly GGC	Tyr UAC	Arg AGG	Gly GGC	Thr	Val GUC	Asn AAU	31 105
319 1056	Thr	Ile	Trp UGG	Asn AAU	Gly GGA	Ile AUU	Pro CCA	Cys UGU	Gln CAG	Arg CGU	Trp UGG	Asp GAU	Ser UCU	Gln CAG	Tyr	Pro -	33 110
335	His	Glu	His	Asp	Met	Thr	Pro	Glu	Asn	Phe	Lys	Cys	Lys	Asp	Leu	Arg	35
1104	CAC	GAG	CAU	GAC	AUG		CCU	GAA	AAU	UUC	AAG	UGC	AAG	GAC	CUA	CGA	115
351	Glu	Asn	Tyr	Cys	Arg	Asn	Pro	Asp	Gly	Ser	Glu	Ser	Pro	Trp	Cys	Phe	36
1152	GAA	AAU	UAC	UGC	CGA	AAU	CCA	GAU	GGG	UCU	GAA	UCA	CCC	UGG	UGU	UUU	1 19
367 1200	Thr	Thr	Asp GAU	Pro	Asn AAC	Ile AUC	Arg CGA	Val GUU	Gly GGC	Tyr UAC	Cys UGC	Ser UCC	Gln CAA	Ile AUU	Pro CCA	Asn AAC	38 124
383	Cys	Asp	Met	Ser	His	Gly	Gln	Asp	Ċys	Tyr	Arg	Gly	nsA	Gly	Lys	Asn	39
1248	UGU	GAU	AUG	UCA	CAU	GGA	CAA	GAU	UGU	UAU	CGU	GGG	UAA	GGC	AAA	AAU	129
399	Tyr	Met	Gly	Asn	Leu	Ser	Gln	Thr	Arg	Ser	Gly	Leu	Thr	Cys	Ser	Met	41
1296	UAU	AUG	GGC	AAC	UUA	UCC	CAA		AGA	UCU	GGA	CUA	ACA	UGU	UCA	AUG	134
415	Trp	Asp	Lys	Asn	Met	Glu	Asp	Leu	His	Arg	His	Ile	Phe	Trp	Glu	Pro	43
1344	UGG	GAC	AAG	AAC	AUG	GAA	GAC	UUA	CAU	CGU	CAU	AUC	UUC	UGG	GAA	CCA	139
431	Asp	Ala	Ser	Lys	Leu	Asn	Glu	Asn	Tyr	Cys	Arg	Asn	Pro	Asp	Asp	Asp	44
1392	GAU	GCA	AGU	AAG	CUG	AAU	GAG	AAU	UAC	UGC	CGA	AAU	CCA	GAU	GAU	GAU	143
447	Ala	His	Gly	Pro	Trp	Cys	Tyr	Thr	Gly	Asn	Pro	Leu	lle	Pro	Trp	Asp	46
1440	GCU	CAU	GGA	CCC	UGG	UGC	UAC	ACG	GGA	AAU	CCA	CUC	AUU	CCU	UGG	GAU	148
463	Tyr	Cys	Pro	Ile	Ser	Arg	Cys	Glu	Gly	Asp	Thr	Thr	Pro	Thr	Ile	Val	47
1488	Uau	UGC	CCU	AUU	UCU	CGU	UGU	GAA	GGU	GAU		ACA	CCU	ACA	AUA	GUC	153
479	Asn	Leu	Asp	His	Pro	Val	Ile	Ser	Cys	Ala	Lys	Thr	Lys	Gln	Leu	Arg	49
1536	AAU	UUA	GAC	CAU	CCC	GUA	AUA	UCU	UGU	GCC	AAA	ACG	AAA	CAA	UUG	CGA	158
495 1584	Val GUU	Val GUA	Asn AAU	Gly GGG	Ile AUU	Pro CCA	Thr	Arg CGA	Thr ACA	Asn AAC	Ile AUA	Gly.	Trp UGG	Met AUG	Val GUU	Ser AGU	51 163
511	Leu	Arg	Tyr	Arg	Asn	Lys	His	Ile	Cys	Gly	Gly	Ser	Leu	Ile	Lys	Glu	52
1632	UUG	AGA	UAC	AGA	AAU	AAA	CAU	AUC	UGC	GGA	GGA	UCA	UUG	AUA	AAG	GAG	157
527	Ser	Trp	Val	Leu	Thr	Ala	Arg	Gln	Cys	Phe	Pro	Ser	Arg	ASP	Leu	Lys	54
1680	AGU	UGG	GUU	CUU	ACU	GCA	CGA	CAG	UGU	UUC	CCU	UCU	CGA	GAC	UUG	AAA	172
543	Asp	Tyr	Glu	Ala	Trp	Leu	Gly	Ile	His	Asp	Val	His	Gly	Arg	Gly	Asp	55
1728	GAU	UAU	GAA	GCU	UGG	CUU	GGA	AUU	CAU	GAU	GUC	CAC	GGA	AGA	GGA	GAU	177
559 1776	Glu	Lys AAA	Cvs	Lys	Gln	Val	Len	4 c n	V-1	C 4 =	C1	•		_			57 182
575 1824	Glu	Gly GGA	Ser	Asp	Leu	Val	Leu	Met	Ive	T 011	415		D			_	59 187

FIG. 12 (cont.)

591 1872	Asp GAU	Asp GAU	Phe UUU	Val GUU	Ser J AGU	Thr	AUU	ASF GAU	Leu J UUA	Pro	RA C	Tyr J UAU	Gly	Cys	Thr	Ile	60 191
607 1920	Pro	Glu	Lvs	Thr	Ser	Cvs	Ser	. V = 1	Tur		· ~		_			Leu UUĞ	62 196
623 1968	Ile	Asn	Tyr	Asn	ดาง	יום ו	T an	4 55 6				_	_			Gly GGA	63
639 2016	Asn	Glu	Lys	Cvs	Ser	Gln	Hie	Hie	450	C1			-	_		Glu GAG	65 206
655 2064	Ser	Glu	Ile	Cvs	Ala	Glv	41.	Glu	1 240	T1-	01-			_		Glu GAG	67. 211
671 2112	Gly	Asp	Tvr	ឲាម	Glv	Pro.	1	3/- 1	0							Met AUG	68 215
687 2160	Val	Leu	Glv	Val	Tle	Va 1	Dwa	Q1			_					Arg CGU	70 220
703 2208	Pro	Gly	Ile	Phe	Val	Ara	Va 1	43-	m			_				Lys AAA	71 225
_716 2256	Ile	Ile	Leu	Thr	Tur	The	Vo 1	D	~ 1	_	_	Leu CUG					73
735 2304	Leu	Lys	His	Pro	Pro	Tla	Gln	1 011	c	D1		*** UGA					230 75
751 2352	Met	Trp	Asn	Leu	T.Ve	Cve	Wie.	T	01-	01	_	*** UAA					235 76
767 2400	Arg	Val	Met	Phe	Va 1	Clu	T1.	7	71.			Tyr					239 78
783 2448	Leu	Phe	Cys	Leu	Ser	Va1	T 011	Dh		۵,	_	*** UGA					244 79
799 2496	Tyr	Met	Gln	Val	***	***	T7 : -	T1 -	_			Tyr					249 81
815 2544	Lys	Lys	His	Thr	Glv	Tle	Phe	A10	000	UGA	AGA	UAC	UUG	UAA	GGA	UUA	254 82
			-110	AUA	uuu	AUA	UUU	GCU	GGA	UGA	UAA						257